

# IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

: U.S. Patent 5,852,195

Issued: December 22, 1998

Inventors : Romines et al.

For : PYRANONE COMPOUNDS USEFUL TO TREAT RETROVIRAL

**INFECTIONS** 

Mail Stop Patent Ext. Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

# APPLICATION FOR EXTENSION OF PATENT TERM UNDER 35 U.S.C. § 156

1

Sir:

Pharmacia & Upjohn Company LLC, a company organized under the laws of Delaware and formerly known as Pharmacia & Upjohn Company (hereinafter called "the Applicant"), is the assignee and owner of record of U.S. Patent 5,852,195 by virtue of an assignment from each of the individual inventors which was recorded on November 23, 1998 at Reel/Frame 009609/0355. The undersigned registered practitioner, acting on behalf of Pharmacia & Upjohn Company LLC, the owner of record, as its attorney, hereby applies for an extension of the term of U.S. Patent 5,852,195 pursuant to 35 U.S.C. § 156 and 37 C.F.R. § 1.710 through § 1.791.

## SUMMARY OF THE APPLICATION FOR EXTENSION

The Applicant seeks extension of the term of U.S. Patent 5,852,195 for a period of 1278 days, so that expiration date of the patent would be changed from 22 December 2015 to 22 June 2019.

This application for patent term extension is predicated upon the approval of an application submitted under section 505(b) of the Federal Food, Drug and Cosmetic Act for the drug APTIVUS® capsules, which was approved by the United States Food and Drug Administration on 22 June 2005 (NDA 21-814).

APTIVUS® capsules is a drug product and its sole active ingredient is tipranavir. Thus, tipranavir is a product which has been subject to a regulatory review period before its commercial marketing or use.

Tipranavir has not been previously approved for commercial marketing or use under the Federal Food, Drug and Cosmetic Act, the Public Health Service Act, or the Virus Serum-Toxin Act.

The patent for which extension is sought claims tipranavir.

Pursuant to 35 U.S.C. § 156(c)(3), the amount of patent term extension is limited to 1278 days.

# DETAILED DESCRIPTION OF THE BASIS FOR THE APPLICATION

The information given below is that which must be included in this application pursuant to 37 C.F.R. § 1.740(a).

# (1) Identification of the Approved Product

The approved product is the compound which is known by the United States Adopted Name (USAN), tipranavir.

Tipranavir has the following structural formula

and is known by the following chemical names:

a) 2-Pyridinesulfonamide, N-[3-[(1R)-1-[(6R)-5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-

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(Preferred CA INDEX NAME);

- b) 2-Pyridinesulfonamide, N-[3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-, [R-(R\*,R\*)]- (Other CA INDEX NAME);
- c) 3'-[(1R)-1-[(6R)-5,6-Dihydro-4-hydroxy-2-oxo-6-phenylethyl-6-propyl-2H-pyran-3yl]propyl]-5-(trifluoromethyl)-2-pyridinesulfonanilide (USP Dictionary of USAN and International Drug Names, 2004 Ed.);
- d) (3R,6R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide; and
- e) 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide.

Tipranavir is also denoted by CAS Registry Number 174484-41-4.

As is evidenced by the text of the labeling (package insert) approved by the U.S. Food and Drug Administration, the approved product, tipranavir, is the active ingredient of APTIVUS® capsules. A copy of the approved labeling is attached hereto as Exhibit A.

# (2) Identification of the Federal Statute Under Which Regulatory Review Occurred

The approved product was the subject of regulatory review under the provisions of Section 505 of the Federal Food, Drug & Cosmetic Act, as amended (21 U.S.C. § 355).

# (3) Date the Product Received Permission for Commercial Marketing

The product received permission for commercial marketing or use under the provisions of Section 505 of the Federal Food, Drug & Cosmetic Act, as amended

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(21 U.S.C. § 355) on 22 June 2005, the date NDA 21-841 was approved by the United States Food and Drug Administration.

# (4) <u>Identification of the Active Ingredient in the Drug Product</u> and Statement that It Has Not Been Previously Approved

APTIVUS® capsules is a drug product. Its sole active ingredient is tipranavir. It is the Applicant's belief that tipranavir has not been previously approved for commercial marketing or use under the Federal Food, Drug, and Cosmetic Act, the Public Health Service Act, or the Virus-Serum-Toxin Act. The use for which tipranavir has been approved is set forth in the label which was approved by the FDA on June 22, 2005 as follows: APTIVUS® (tipranavir), co-administered with 200 mg of ritonavir, is indicated for combination antiretroviral treatment of HIV-1 infected adult patients with evidence of viral replication, who are highly treatment-experienced or have HIV-1 strains resistant to multiple protease inhibitors. The provision of law under which the commercial marketing or use of tipranavir was approved is Section 505 of the Federal Food, Drug & Cosmetic Act, as amended (21 U.S.C. § 355).

# (5) Application is Being Submitted Within Sixty Day Period Permitted

This application is being submitted within the sixty day period permitted for submission pursuant to § 1.720(f). Such sixty day period will expire on 21 August 2005, but because that date falls on a Sunday, the last day on which the application for patent term extension could be submitted is 22 August 2005, pursuant to 37 C.F.R. § 1.7.

# (6) Identification of the Patent for Which Extension is Sought

The patent for which an extension is being sought is U.S. Patent 5,852,195 (hereinafter referred to as the "'195 patent').

The inventors named in the patent are ROMINES, KAREN RENE; BUNDY, GORDON L; SCHWARTZ, THERESA M.; TOMMASI, RUBEN A.; STROHBACH, JOSEPH W.; TURNER, STEVEN, RONALD; THAISRIVONGS, SUVIT; ARISTOFF, PAUL ADRIAN; JOHNSON, PAUL D; SKULNICK, HARVEY IRVING; SKALETZKY, LOUIS L.; ANDERSON, DAVID JOHN; MORRIS, JOEL; GAMMILL, RONALD B.; and LUKE, GEORGE P.

The '195 patent issued on 22 December 1998.

Pursuant to 35 U.S.C. § 154(c), and absent any extension, the term of the '195 patent will expire on 22 December 2015.

# (7) Copy of the Patent for Which Extension is Sought

A copy of the '195 patent, including the entire specification (including claims) and drawings is attached hereto as Exhibit B.

# (8) Copy of any Disclaimer, Certificate of Correction, Receipt of Maintenance Fee Payment, or Reexamination Certificate Issued in the Patent

No disclaimers have been issued for the '195 patent.

A copy of the certificate of correction issued on 9 October 2001 for the '195 is attached hereto as Exhibit C. No other certificates of correction have been issued.

A copy of a maintenance fee statement showing that the first maintenance fee has been paid is attached hereto as Exhibit D. The second maintenance fee is not yet due.

The '195 patent has not been subjected to reexamination.

# (9) Statement that the Patent Claims the Approved Product and Showing that Claims Read on the Approved Product

The sole active ingredient of the drug product APTIVUS® capsules is tipranavir. As noted previously, tipranavir has the following structural formula

and is known by, inter alia, the chemical name (3R,6R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H- pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide.

The '195 patent contains 5 claims. Claim 1, the sole independent claim, reads as follows:

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# 1. The compound of the formula VI

wherein R<sub>2</sub> is

- a)  $H_3C$ — $CH_2$ —, or
- b) phenyl- $(CH_2)_2$ —;

wherein R<sub>3</sub> is the moiety of formula X

wherein Ro is

- a)  $H_3C (CH_2)_2$ , or
  - b) phenyl- $(CH_2)_2$ —;

wherein R<sub>7</sub> is H<sub>3</sub>C-CH<sub>2</sub>-;

wherein Ro is -NHSO2-het;

wherein het is 2-pyridinyl substituted at the 5-position by zero (0) or one (1)  $R_{10}$ ;

wherein R<sub>10</sub> is

- a) —CN,
- b) —CF<sub>3</sub>,
- c) -NH2, or
- d) --CONH,;

or a pharmaceutically acceptable salt thereof.

The certificate of correction dated October 9, 2001, a copy of which is included herewith as Exhibit C, corrects claim 1 by replacing " $R_y$ " with " $R_7$ ".

Claim 1 reads on the approved product because tipranavir is the compound of the formula

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wherein,

 $R_2$  is phenyl-(CH<sub>2</sub>)<sub>2</sub>-;

R<sub>3</sub> is the moiety of formula

wherein,

R<sub>7</sub> is H<sub>3</sub>C-CH<sub>2</sub>-,

R<sub>9</sub> is -NHSO<sub>2</sub>-het,

het is 2-pyridinyl substituted at the 5-position by R<sub>10</sub>, and

R<sub>10</sub> is-CF<sub>3</sub>; and

wherein  $R_6$  is  $H_3C$ -(CH2)<sub>2</sub>-.

Claim 3 reads on the approved product because the fourth compound of the Markush listing of claim 3 corresponds to the species tiparanvir as well to as all stereoisomers of tipranavir, and because the fifth compound of the Markush listing of claim 3 is the species tipranavir.

Claim 4 reads on the approved product because the first compound of the Markush listing of claim 4 is tipranavir.

# (10) Information to Enable Determination of the Regulatory Review Period

The relevant dates and information needed to enable the Secretary of Health and Human Services to determine the applicable regulatory review period appears in Exhibit E, which is attached hereto.

# (11) Significant Activities by the Marketing Applicant during the Regulatory Review Period

A description of the significant activities undertaken by the marketing applicant during the applicable regulatory review period with respect to the approved product and the significant dates applicable to such activities is provided, on a separate page, by the attached Exhibit F. This exhibit includes four chronologies:

- Agency Contact Reports Any phone calls, meetings or emails between BI and FDA
- Correspondence from FDA Faxes or letters (hard copy) from FDA to BI
- IND Log Listing of all submissions to the tipranavir IND 51,979
- NDA Log Listing of all amendments to the tipranavir NDA 21-814 (NDA 21-822 (solution) cross references 21-814 for all clinical and non-clinical data)

# (12) Statement of Opinion of the Applicant that the Patent is Eligible for Extension Claimed

The attached Exhibit G provides the required statement, on a separate page, that in the opinion of the applicant the patent is eligible for the extension and a statement as to the length of extension claimed, including how the length of extension was determined.

# (13) Statement Acknowledging Duty Of Disclosure

Applicant acknowledges a duty to disclose to the Director of the United States Patent and Trademark Office and the Secretary of Health and Human Services any information which is material to the determination of entitlement to the extension sought.

# (14) Payment of Prescribed Fee

The Commissioner is authorized to charge the fee prescribed by 37 C.F.R. § 1.20(j) for receiving and acting upon this application for extension (\$1,120.00) to Deposit Account No. 02-2955 (Deposit Account Name: Boehringer Ingelheim Corporation). The Commissioner is authorized to charge any additional fees or underpayments due for consideration of this application, and to credit any overpayments, to this same deposit account.

# (15) Correspondence

Correspondence relating to this application for patent term extension should be directed to:

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A total of five copies of this application are being submitted, in accordance with MPEP Section 2753.

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# **SUMMATION**

Having included in this application all of the requisite information required, the Applicant requests extension of the term of U.S. Patent 5,852,195 for a period of 1278 days.

Respectfully submitted,

Alan Stempel

Attorney for the Applicant

Reg. No. 28,991

Date: August 16, 2005

List of Enclosures / Attachments

- (1) Return Post Card
- (2) Exhibits A-H

EXHIBIT A COPY OF APPROVED LABELING FOR APTIVUS® CAPSULES

ATTENTION PHARMACIST: Detach "Patient's Instructions for Use" from package insert and dispense with product. Dispense the capsules in the unit of use container.

# **Aptivus®**

(tipranavir)

# Capsules, 250 mg



#### **Prescribing Information**

## WARNING

APTIVUS CO-ADMINISTERED WITH 200 MG RITONAVIR HAS BEEN ASSOCIATED WITH REPORTS OF CLINICAL HEPATITIS AND HEPATIC DECOMPENSATION INCLUDING SOME FATALITIES. EXTRA VIGILANCE IS WARRANTED IN PATIENTS WITH CHRONIC HEPATITIS B OR HEPATITIS C CO-INFECTION, AS THESE PATIENTS HAVE AN INCREASED RISK OF HEPATOTOXICITY. (SEE WARNINGS)

#### DESCRIPTION

APTIVUS® (tipranavir) is the brand name for tipranavir (TPV), a non-peptidic protease inhibitor (PI) of HIV belonging to the class of 4-hydroxy-5,6-dihydro-2-pyrone sulfonamides.

APTIVUS soft gelatin capsules are for oral administration. Each capsule contains 250 mg tipranavir. The major inactive ingredients in the capsule are dehydrated alcohol (7% w/w or 0.1 g per capsule), polyoxyl 35 castor oil, propylene glycol, mono/diglycerides of caprylic/capric acid and gelatin.

The chemical name of tipranavir is 2-Pyridinesulfonamide, N-[3-[(1R)-1-[(6R)-5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl). It has a molecular formula of  $C_{31}H_{33}F_3N_2O_5S$  and a molecular weight of 602.7. Tipranavir has the following structural formula and is a single stereoisomer with the 1R, 6R configuration.

Tipranavir is a white to off-white to slightly yellow solid. It is freely soluble in dehydrated alcohol and propylene glycol, and insoluble in aqueous buffer at pH 7.5.

#### **CLINICAL PHARMACOLOGY**

# Microbiology

#### Mechanism of Action

Tipranavir (TPV) is a non-peptidic HIV-1 protease inhibitor that inhibits the virus-specific processing of the viral Gag and Gag-Pol polyproteins in HIV-1 infected cells, thus preventing formation of mature virions.

# Antiviral Activity

Tipranavir inhibits the replication of laboratory strains of HIV-1 and clinical isolates in acute models of T-cell infection, with 50% effective concentrations (EC<sub>50</sub>) ranging from 0.03 to 0.07 μM (18-42 ng/mL). Tipranavir demonstrates antiviral activity *in vitro* against a broad panel of HIV-1 group M non-clade B isolates (A, C, D, F, G, H, CRF01 AE, CRF02 AG, CRF12 BF). Group O and HIV-2 isolates have reduced susceptibility *in vitro* to tipranavir with EC<sub>50</sub> values ranging from 0.164 -1 μM and 0.233-0.522 μM, respectively. Protein binding studies have shown that the antiviral activity of tipranavir decreases on average 3.75-fold in conditions where human serum is present. When used with other antiretroviral agents *in vitro*, the combination of tipranavir was additive to antagonistic with other protease inhibitors (amprenavir, atazanavir, indinavir, lopinavir, nelfinavir, ritonavir, and saquinavir) and generally additive with the NNRTIs (delavirdine, efavirenz, and nevirapine) and the NRTIs (abacavir, didanosine, emtricitabine, lamivudine, stavudine, tenofovir, and zidovudine). Tipranavir was synergistic with the HIV fusion inhibitor enfuvirtide. There was no antagonism of the *in vitro* combinations of tipranavir with either adefovir or ribavirin, used in the treatment of viral hepatitis.

#### Resistance

In vitro: HIV-1 isolates with a decreased susceptibility to tipranavir have been selected in vitro and obtained from patients treated with APTIVUS/ritonavir (TPV/ritonavir). HIV-1 isolates that were 87-fold resistant to tipranavir were selected in vitro by 9 months and contained 10 protease mutations that developed in the following order: L33F, I84V, K45I, I13V, V32I, V82L, M36I, A71V, L10F, and I54V/T. Changes in the Gag polyprotein CA/P2 cleavage site were also observed following drug selection. Experiments with site-directed mutants of HIV-1 showed that the presence of 6 mutations in the protease coding sequence (I13V, V32I, L33F, K45I, V82L, I84V) conferred > 10-fold reduced susceptibility to tipranavir. Recombinant viruses showing ≥ 3-fold reduced susceptibility to tipranavir were growth impaired.

Clinical Studies of Treatment-Experienced Patients: In Phase 3 studies 1182.12 and 1182.48, multiple protease inhibitor-resistant HIV-1 isolates from 59 highly treatment-experienced patients who received APTIVUS/ritonavir and experienced virologic rebound developed amino acid substitutions that were associated with resistance to tipranavir. The most common amino acid substitutions that developed on 500/200mg APTIVUS/ritonavir in greater than 20% of APTIVUS/ritonavir virologic failure isolates were L33V/I/F, V82T, and I84V. Other substitutions that developed in 10 to 20% of APTIVUS/ritonavir virologic failure isolates included L10V/I/S, I13V, E35D/G/N, I47V, K55R, V82L, and L89V/M. Tipranavir resistance was detected at virologic rebound after an average of

38 weeks of APTIVUS/ritonavir treatment with a median 14-fold decrease in tipranavir susceptibility. The resistance profile in treatment-naïve subjects has not been characterized.

#### Cross-resistance

Cross-resistance among protease inibitors has been observed. Tipranavir had < 4-fold decreased susceptibility against 90% (94/105) of HIV-1 isolates resistant to amprenavir, atazanavir, indinavir, lopinavir, nelfinavir, ritonavir, or saquinavir. Tipranavir-resistant viruses which emerged *in vitro* had decreased susceptibility to the protease inhibitors amprenavir, atazanavir, indinavir, lopinavir, nelfinavir and ritonavir but remained sensitive to saquinavir.

# **Baseline Genotype and Virologic Outcome Analyses**

Genotypic and/or phenotypic analysis of baseline virus may aid in determining tipranavir susceptibility before initiation of APTIVUS/ritonavir therapy. Several analyses were conducted to evaluate the impact of specific mutations and mutational patterns on virologic outcome. Both the type and number of baseline protease inhibitor mutations as well as use of additional active agents (e.g., enfuvirtide) affected APTIVUS/ritonavir response rates in Phase 3 studies 1182.12 and 1182.48 through Week 24 of treatment.

Regression analyses of baseline and/or on-treatment HIV-1 genotypes from 860 highly treatment-experienced patients in Phase 2 and 3 studies demonstrated that mutations at 16 amino acid codons in the HIV protease coding sequence were associated with reduced virologic responses at 24 weeks and/or reduced tipranavir susceptibility: L10V, I13V, K20M/R/V, L33F, E35G, M36I, K43T, M46L, I47V, I54A/M/V, Q58E, H69K, T74P, V82L/T, N83D or I84V.

Analyses were also conducted to assess virologic outcome by the number of primary protease inhibitor mutations present at baseline. Response rates were reduced if five or more protease inhibitor-associated mutations were present at baseline and subjects did not receive concomitant enfuvirtide with APTIVUS/ritonavir. See Table 1.

Table 1 Phase 3 Studies 1182.12 and 1182.48: Proportion of Responders (confirmed  $\geq 1 \log_{10}$  decrease at Week 24) by Number of Baseline Primary Protease Inhibitor (PI) Mutations

Number of Baseline Primary PI Mutations <sup>a</sup>	APTIVUS/ritonavir N = 513		Comparator PI/ritonavir N = 502		
	No Enfuvirtide	+ Enfuvirtide	No Enfuvirtide	+ Enfuvirtide	
Overall	40% (147/368)	<b>64%</b> (93/145)	19% (75/390)	<b>30%</b> (34/112)	
1 - 2	68% (26/38)	75% (3/4)	41% (17/41)	100% (2/2)	
3 - 4	44% (78/176)	<b>64%</b> (39/61)	23% (39/170)	<b>40%</b> (21/52)	
5+	<b>28%</b> (43/151)	<b>64%</b> (51/80)	11% (19/178)	19% (11/57)	

<sup>&</sup>lt;sup>a</sup> Primary PI mutations include any amino acid change at positions 30, 32, 36, 46, 47, 48, 50, 53, 54, 82, 84, 88 and 90

The median change from baseline in HIV-1 RNA at weeks 2, 4, 8, 16 and 24 was evaluated by the number of baseline primary protease inhibitor mutations (1-4 or  $\geq$  5) in subjects who received APTIVUS/ritonavir with or without enfuvirtide. The following observations were made:

- Approximately 1.5 log<sub>10</sub> decrease in HIV-1 RNA at early time points (Week 2) regardless of the number of baseline primary protease inhibitor mutations (1-4 or 5+).
- Subjects with 5 or more primary protease inhibitor mutations in their HIV-1 at baseline who received APTIVUS/ritonavir without enfuvirtide (n=204) began to lose their antiviral response after Week 4.
- Early HIV-1 RNA decreases (1.5-2 log<sub>10</sub>) were sustained through Week 24 in subjects with 5 or more primary protease inhibitor mutations at baseline who received enfuvirtide with APTIVUS/ritonavir (n=88).

Conclusions regarding the relevance of particular mutations or mutational patterns are subject to change pending additional data.

# **Baseline Phenotype and Virologic Outcome Analyses**

APTIVUS/ritonavir response rates were also assessed by baseline tipranavir phenotype. Relationships between baseline phenotypic susceptibility to tipranavir, mutations at protease amino acid codons 33, 82, 84 and 90, tipranavir resistance-associated mutations, and response to APTIVUS/ritonavir therapy at Week 24 are summarized in Table 2. These baseline phenotype groups are not meant to represent clinical susceptibility breakpoints for APTIVUS/ritonavir because the data are based on the select 1182.12 and 1182.48 patient population. The data are provided to give clinicians information on the likelihood of virologic success based on pre-treatment susceptibility to APTIVUS/ritonavir in highly protease inhibitor-experienced patients.

Table 2 Response by Baseline Tipranavir Phenotype in the 1182.12 and 1182.48 Trials

Baseline Tipranavir Phenotype (Fold Change) <sup>a</sup>	Proportion of Responders with No Enfuvirtide Use	Proportion of Responders <sup>b</sup> with ENF Use	# of Baseline Protease Mutations at 33, 82, 84, 90	# of Baseline Tipranavir Resistance- Associated Mutations <sup>c</sup>	Tipranavir Susceptibility
0-3	45% (74/163)	77% (46/60)	0-2	0-4	Susceptible
> 3-10	21% (10/47)	43% (12/28)	3	5-7	Decreased Susceptibility
> 10	0% (0/8)	57% (4/7)	4	8+	Resistant

<sup>&</sup>lt;sup>a</sup>Change in tipranavir IC<sub>50</sub> value from wild-type reference

<sup>&</sup>lt;sup>o</sup>Confirmed ≥1 log<sub>10</sub> decrease at Week 24

<sup>&</sup>lt;sup>c</sup>Number of amino acid substitutions in HIV protease among L10V, I13V, K20M/R/V, L33F, E35G, M36I, K43T, M46L, I47V, I54A/M/V, Q58E, H69K, T74P, V82L/T, N83D or I84V

# **Pharmacodynamics**

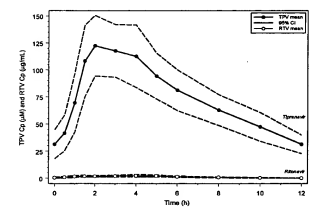
The median Inhibitory Quotient (IQ) determined from 301 highly treatment-experienced patients was about 75 (inter-quartile range: 29-189), from pivotal clinical trials 1182.12 and 1182.48. The IQ is defined as the tipranavir trough concentration divided by the viral IC<sub>50</sub> value, corrected for protein binding. There was a relationship between the proportion of patients with a  $\geq$  1 log<sub>10</sub> reduction of viral load from baseline at week 24 and their IQ value. Among the 206 patients receiving APTIVUS/ritonavir without enfuvirtide, the response rate was 23% in those with an IQ value < 75 and 55% in those with an IQ value  $\geq$  75. Among the 95 patients receiving APTIVUS/ritonavir with enfuvirtide, the response rates in patients with an IQ value < 75 versus those with an IQ value  $\geq$  75 were 43% and 84%, respectively. These IQ groups are derived from a select population and are not meant to represent clinical breakpoints.

#### **Pharmacokinetics in Adult Patients**

In order to achieve effective tipranavir plasma concentrations and a twice-daily dosing regimen, co-administration of APTIVUS with 200 mg of ritonavir is essential (see PRECAUTIONS and DOSAGE AND ADMINISTRATION). Ritonavir inhibits hepatic cytochrome P450 3A (CYP 3A), the intestinal P-glycoprotein (P-gp) efflux pump and possibly intestinal CYP 3A. In a dose-ranging evaluation in 113 HIV-negative male and female volunteers, there was a 29-fold increase in the geometric mean morning steady-state trough plasma concentrations of tipranavir following tipranavir co-administered with low-dose ritonavir (500/200 mg twice daily) compared to tipranavir 500 mg twice daily without ritonavir.

Figure 1 displays mean plasma concentrations of tipranavir and ritonavir at steady state for the 500/200 mg tipranavir/ritonavir dose.

Figure 1 Mean Steady State Tipranavir Plasma Concentrations (95% CI) with Ritonavir Co-administration (tipranavir/ritonavir 500/200 mg BID)



# Absorption and Bioavailability

Absorption of tipranavir in humans is limited, although no absolute quantification of absorption is available. Tipranavir is a P-gp substrate, a weak P-gp inhibitor, and appears to be a potent P-gp inducer as well. *In vivo* data suggest that the net effect of tipranavir/ritonavir at the proposed dose regimen (500/200 mg) is P-gp induction at steady-state, although ritonavir is a P-gp inhibitor. Tipranavir trough concentrations at steady-state are about 70% lower than those on Day 1, presumably due to intestinal P-gp induction. Steady state is attained in most subjects after 7-10 days of dosing.

Dosing with APTIVUS 500 mg concomitant with 200 mg ritonavir twice-daily for greater than 2 weeks and without meal restriction produced the following pharmacokinetic parameters for female and male HIV-positive patients. See Table 3.

Table 3 Pharmacokinetic Parameters<sup>a</sup> of tipranavir/ritonavir 500/200 mg for HIV+ Patients by Gender

	Females (n = 14)	Males (n = 106)
p <sub>trough</sub> (μM)	$41.6 \pm 24.3$	$35.6 \pm 16.7$
C <sub>max</sub> (μM)	$94.8 \pm 22.8$	$77.6 \pm 16.6$
max (h)	2.9	3.0
UC <sub>0-12h</sub> (μM•h)	851 ± 309	710 ± 207
L (L/h)	1.15	1.27
/ (L)	7.7	10.2
1/2 (h)	5.5	6.0

<sup>&</sup>lt;sup>a</sup>Population pharmacokinetic parameters reported as mean ± standard deviation

## **Effects of Food on Oral Absorption**

APTIVUS capsules co-administered with ritonavir should be taken with food. Bioavailability is increased with a high fat meal. Tipranavir capsules, administered under high fat meal conditions or with a light snack of toast and skimmed milk, were tested in a multiple dose study. High-fat meals (868 kcal, 53% derived from fat, 31% derived from carbohydrates) enhanced the extent of bioavailability (AUC point estimate 1.31, confidence interval 1.23-1.39), but had minimal effect on peak tipranavir concentrations ( $C_{max}$  point estimate 1.16, confidence interval 1.09-1.24).

When APTIVUS, co-administered with low-dose ritonavir, was co-administered with 20 mL of aluminum and magnesium-based liquid antacid, tipranavir AUC<sub>12h</sub>, C<sub>max</sub> and C<sub>12h</sub> were reduced by 25-29%. Consideration should be given to separating tipranavir/ritonavir dosing from antacid administration to prevent reduced absorption of tipranavir.

#### Distribution

Tipranavir is extensively bound to plasma proteins (> 99.9%). It binds to both human serum albumin and  $\alpha$ -1-acid glycoprotein. The mean fraction of APTIVUS (dosed without ritonavir) unbound in plasma was similar in clinical samples from healthy volunteers (0.015%  $\pm$  0.006%) and HIV-positive patients (0.019%  $\pm$  0.076%). Total plasma tipranavir concentrations for these samples ranged from 9 to 82  $\mu$ M. The unbound fraction of tipranavir appeared to be independent of total drug concentration over this concentration range.

No studies have been conducted to determine the distribution of tipranavir into human cerebrospinal fluid or semen.

#### Metabolism

*In vitro* metabolism studies with human liver microsomes indicated that CYP 3A4 is the predominant CYP enzyme involved in tipranavir metabolism.

The oral clearance of tipranavir decreased after the addition of ritonavir, which may represent diminished first-pass clearance of the drug at the gastrointestinal tract as well as the liver.

The metabolism of tipranavir in the presence of 200 mg ritonavir is minimal. Administration of <sup>14</sup>C-tipranavir to subjects that received tipranavir/ritonavir 500/200 mg dosed to steady-state demonstrated that unchanged tipranavir accounted for 98.4% or greater of the total plasma radioactivity circulating at 3, 8, or 12 hours after dosing. Only a few metabolites were found in plasma, and all were at trace levels (0.2% or less of the plasma radioactivity). In feces, unchanged tipranavir represented the majority of fecal radioactivity (79.9% of fecal radioactivity). The most abundant fecal metabolite, at 4.9% of fecal radioactivity (3.2% of dose), was a hydroxyl metabolite of tipranavir. In urine, unchanged tipranavir was found in trace amounts (0.5% of urine radioactivity). The most abundant urinary metabolite, at 11.0% of urine radioactivity (0.5% of dose) was a glucuronide conjugate of tipranavir.

## Elimination

Administration of <sup>14</sup>C-tipranavir to subjects (n=8) that received tipranavir/ritonavir 500/200 mg dosed to steady-state demonstrated that most radioactivity (median 82.3%) was excreted in feces, while only a median of 4.4% of the radioactive dose administered was recovered in urine. In addition, most radioactivity (56%) was excreted between 24 and 96 hours after dosing. The effective mean elimination half-life of tipranavir/ritonavir in healthy volunteers (n=67) and HIV-infected adult patients (n=120) was approximately 4.8 and 6.0 hours, respectively, at steady state following a dose of 500/200 mg twice daily with a light meal.

# Pharmacokinetics in Special Populations

#### Renal Impairment

APTIVUS pharmacokinetics has not been studied in patients with renal dysfunction. However, since the renal clearance of tipranavir is negligible, a decrease in total body clearance is not expected in patients with renal insufficiency.

# Hepatic Impairment

In a study comparing 9 patients with mild (Child-Pugh A) hepatic impairment to 9 controls, the single and multiple dose plasma concentrations of tipranavir and ritonavir were increased in patients with hepatic impairment, but were within the range observed in clinical trials. No dosing adjustment is required in patients with mild hepatic impairment.

The influence of moderate hepatic impairment (Child-Pugh B) or severe hepatic impairment (Child-Pugh C) on the multiple-dose pharmacokinetics of tipranavir administered with ritonavir has not been evaluated (see DOSAGE AND ADMINISTRATION, CONTRAINDICATIONS, and WARNINGS).

#### Gender

Evaluation of steady-state plasma tipranavir trough concentrations at 10-14 h after dosing from the 1182.12 and 1182.48 studies demonstrated that females generally had higher tipranavir concentrations than males. After 4 weeks of tipranavir/ritonavir 500/200 mg BID, the median plasma trough concentration of tipranavir was 43.9  $\mu$ M for females and 31.1  $\mu$ M for males. The difference in concentrations does not warrant a dose adjustment.

#### Race

Evaluation of steady-state plasma tipranavir trough concentrations at 10-14 h after dosing from the 1182.12 and 1182.48 studies demonstrated that white males generally had more variability in tipranavir concentrations than black males, but the median concentration and the range making up the majority of the data are comparable between the races.

## Geriatric Patients

Evaluation of steady-state plasma tipranavir trough concentrations at 10-14 h after dosing from the 1182.12 and 1182.48 studies demonstrated that there was no change in median trough tipranavir concentrations as age increased for either gender through 65 years of age. There were an insufficient number of women greater than age 65 years in the two trials to evaluate the elderly, but the trend of consistent trough tipranavir concentrations with increasing age through 80 years for men was supported.

#### Pediatric Patients

The pharmacokinetic profile of tipranavir in pediatric patients has not been established.

## **Drug Interactions**

See also CONTRAINDICATIONS, WARNINGS and PRECAUTIONS, Drug Interactions.

APTIVUS co-administered with 200 mg of ritonavir can alter plasma exposure of other drugs and other drugs may alter plasma exposure of tipranavir.

# Potential for tipranavir/ritonavir to Affect Other Drugs

- 1. APTIVUS co-administered with 200 mg of ritonavir at the recommended dose, is a net inhibitor of CYP 3A and may increase plasma concentrations of agents that are primarily metabolized by CYP 3A. Thus, co-administration of APTIVUS/ritonavir with drugs highly dependent on CYP 3A for clearance and for which elevated plasma concentrations are associated with serious and/or life-threatening events should be contraindicated. Co-administration with other CYP 3A substrates may require a dose adjustment or additional monitoring (see CONTRAINDICATIONS and PRECAUTIONS).
- 2. Studies in human liver microsomes indicated tipranavir is an inhibitor of CYP 1A2, CYP 2C9, CYP 2C19 and CYP 2D6. The potential net effect of tipranavir/ritonavir on CYP 2D6 is inhibition, because ritonavir is a CYP 2D6 inhibitor. The *in vivo* net effect of tipranavir administered with ritonavir on CYP 1A2, CYP 2C9 and CYP 2C19 is not known. Data are not available to indicate whether tipranavir inhibits or induces glucuronosyl transferases and whether tipranavir induces CYP 1A2, CYP 2C9 and CYP 2C19.
- 3. Tipranavir is a P-gp substrate, a weak P-gp inhibitor, and appears to be a potent P-gp inducer as well. Data suggest that the net effect of tipranavir co-administered with 200 mg of ritonavir is P-gp induction at steady-state, although ritonavir is a P-gp inhibitor.
- 4. It is difficult to predict the net effect of APTIVUS administered with ritonavir on oral bioavailability and plasma concentrations of drugs that are dual substrates of CYP 3A and P-gp. The net effect will vary depending on the relative affinity of the co-administered drugs for CYP 3A and P-gp, and the extent of intestinal first-pass metabolism/efflux.

## Potential for Other Drugs to Affect tipranavir

- 1. Tipranavir is a CYP 3A substrate and a P-gp substrate. Co-administration of APTIVUS/ritonavir and drugs that induce CYP 3A and/or P-gp may decrease tipranavir plasma concentrations. Co-administration of APTIVUS/ritonavir and drugs that inhibit P-gp may increase tipranavir plasma concentrations.
- 2. Co-administration of APTIVUS/ritonavir with drugs that inhibit CYP 3A may not further increase tipranavir plasma concentrations, because the level of metabolites is low following steady-state administration of APTIVUS/ritonavir 500/200 mg twice daily.

Drug interaction studies were performed with APTIVUS, co-administered with 200 mg of ritonavir, and other drugs likely to be co-administered and some drugs commonly used as probes for pharmacokinetic interactions. The effects of co-administration of APTIVUS with 200 mg ritonavir, on the AUC,  $C_{max}$  and  $C_{min}$ , are summarized in Tables 4 and 5. For information regarding clinical recommendations (see **PRECAUTIONS**, **Drug Interactions**, **Tables 8** and **9**).

Table 4 Drug Interactions: Pharmacokinetic Parameters for Tipranavir in the Presence of Co-administered Drugs

Co- administered Drug	Co- administered Drug Dose	TPV/ritonavir Drug Dose (Schedule)	n	PK	Pharmacok Co	onfidence Interval) o inetic Parameters wit o-administered Drug No Effect = 1.00	th/without ;
Ü	(Schedule)	` ,			C <sub>max</sub>	AUC	C <sub>min</sub>
Atorvastatin	10 mg (1 dose)	500/200 mg BID (14 doses)	22	↔	0.96 (0.86, 1.07)	1.08 (1.00, 1.15)	1.04 (0.89, 1.22)
Clarithromycin	500 mg BID (25 doses)	500/200 mg BID*	24(68)	<b>↑</b>	1.40 (1.24, 1.47)	1.66 (1.43, 1.73)	2.00 (1.58, 2.47)
Didanosine	400 mg (1 dose)	500/100 mg BID (27 doses)	5	Ţ	1.32 (1.09, 1.60)	1.08 (0.82, 1.42)	0.66 (0.31, 1.43)
Efavirenz	600 mg QD (8 doses)	500/100 mg BID*	21(89)	1	0.79 (0.69, 0.89)	0.69 (0.57, 0.83)	0.58 (0.36, 0.86)
		750/200 mg BID*	25(100)	$\leftrightarrow$	0.97 (0.85,1.09)	1.01 (0.85 , 1.18)	0.97 (0.69 , 1.28)
Ethinyl estradiol /Norethindrone	0.035/1.0 mg (1 dose)	500/100 mg BID (21 doses)	21	1	1.10 (0.98, 1.24)	0.98 (0.88, 1.11)	0.73 (0.59, 0.90)
		750/200 mg BID (21 doses)	13	$\leftrightarrow$	1.01 (0.96, 1.06)	0.98 (0.90, 1.07)	0.91 (0.69, 1.20)
Fluconazole	100 mg QD (12 dose)	500/200 mg BID*	20(68)	1	1.32 (1.18 ,1.47)	1.50 (1.29 , 1.73)	1.69 (1.33, 2.09)
Loperamide	16 mg (1 dose)	750/200 mg BID (21 doses)	24	<b>\</b>	1.03 (0.92, 1.17)	0.98 (0.86, 1.12)	0.74 (0.62, 0.88)
Rifabutin	150 mg (1 dose)	500/200 mg BID (15 doses)	21	$\leftrightarrow$	0.99 (0.93, 1.07)	1.00 (0.96, 1.04)	1.16 (1.07, 1.27)
Tenofovir	300 mg (1 dose)	500/100 mg BID	22	1	0.83 (0.74, 0.94)	0.82 (0.75, 0.91)	0.79 (0.70, 0.90)
		750/200 mg BID (23 doses)	20	$\leftrightarrow$	0.89 (0.84, 0.96)	0.91 (0.85, 0.97)	0.88 (0.78, 1.00)
Zidovudine	300 mg (1 dose)	500/100 mg BID	29	1	0.87 (0.80, 0.94)	0.82 (0.76, 0.89)	0.77 (0.68, 0.87)
	, ,	750/200 mg BID (23 doses)	25	$\leftrightarrow$	1.02 (0.94, 1.10)	1.02 (0.92, 1.13)	1.07 (0.86, 1.34)

<sup>\*</sup>steady state comparison to historical data

Table 5 Drug Interactions: Pharmacokinetic Parameters for Co-administered Drug in the Presence of tipranavir/ritonavir

Clear   Cle	Co-administered Drug	Co- administered Drug Dose	TPV/ritonavir Drug Dose	n	PK	Drug Pharma	nfidence Interval) of acokinetic Paramete TPV/ritonavir; No Effect = 1.00	rs with/without
Amprenavir/RTV b         600/100 mg BID (27 doses)         500/200 mg BID (28 doses)         16 ↓ 1 ↓ 0.61 (0.51, 0.73) do.56 (0.49, 0.64) do.45 (0.38, 0.53) do.44 (0.39, 0.49) so.44 (0.39, 0.49) do.47 (0.39, 0.49) so.44 (0.39, 0.49) do.47 (0.63) do.44 (0.35, 0.74) do.47 (0.63) do.44 (0.55, 0.74) do.48 (0.42, 0.53) do.56 (0.55, 0.76) do.47 (0.64) do.47 (0.63) do.44 (0.55, 0.74) do.48 (0.42, 0.53) do.56 (0.55, 0.76) do.47 (0.64) do.47 (0.63) do.44 (0.47, 0.63) do.45 (0.47, 0.63) do.44 (0.47, 0.63) do.45 (0.47, 0.63) do.44 (0.47, 0.63) do.44 (0.47, 0.63) do.45 (0.47, 0.63) do.44 (0.47, 0.63) do.		(Schedule)	(Schedule)			Cmax	AUC	Cmin
Abacavir a 300 mg BID	Amprenavir/RTV a	600/100 mg BID				ď	0.56 (0.49, 0.64)	d
Abacavir	Ampienaviinerv	(27 doses)	(28 doses)	74	1	-	-	
Adactivation   Adactivation   Adactivation   Adaptivation   Ada	A boossis 8	300 mg BID	250/200 mg BID	28	1	0.56 (0.48, 0.66)	0.56 (0.49, 0.63)	•
Atorvastatin 10 mg (1 dose)	Abacavir		750/100 mg BID	14	1	0.54 (0.47, 0.63)	0.64 (0.55, 0.74)	-
Orthohydroxy-atorivastatin  Orthohydroxy-atorivastatin  Parahydroxy-atorivastatin  Parahydroxy-atorivastatin  Parahydroxy-atorivastatin  Parahydroxy-atorivastatin  Parahydroxy-atorivastatin  Orthohydroxy-atorivastatin  Parahydroxy-atorivastatin  Orthohydroxy-atorivastatin  Parahydroxy-atorivastatin  Orthohydroxy-atorivastatin  Orthohyd				11	Ţ	0.48 (0.42, 0.53)	0.65 (0.55, 0.76)	-
Parahydroxy-atorivastatin    12	Atorvastatin			22	1	8.61 (7.25, 10.21)	9.36 (8.02, 10.94)	5.19 (4.21, 6.40)
Parahydroxy-atorvastatin	Orthohydroxy-ato		, ,	12,	Ţ	0.02 (0.02, 0.03)	0.11 (0.08, 0.17)	0.07 (0.06, 0.08)
14-OH-clarithromycin   21	Parahydroxy-ator	vastatin		13, 22,	<b>\</b>	1.04 (0.87, 1.25)	0.18 (0.14, 0.24)	0.33 (NA)
Didanosine    200 mg BID,   250/200 mg BID   10	Clarithromycin			21	1	0.95 (0.83, 1.09)	1.19 (1.04, 1.37)	1.68 (1.42, 1.98)
Didanosine   200 fig BID   250/200 mg BID   8	14-OH-clarithron	nycin	,	21	Ţ	0.03 (0.02, 0.04)	0.03 (0.02, 0.04)	0.05 (0.04, 0.07)
250 kg	Didonosino	200 mg BID,	250/200 mg BID	10	$\overline{}$	0.57 (0.42, 0.79)	0.67 (0.51, 0.88)	-
125 mg BID,	Diganosine	≥60 kg	750/100 mg BID	8	$\leftrightarrow$		0.97 (0.64, 1.47)	-
A00 mg (1 dose)		<60 kg		9	$\leftrightarrow$	0.77 (0.47,1.26)	0.87 (0.47, 1.65)	-
Efavirenz   Company   C			500/100 mg BID			0.80 (0.63, 1.02)	0.90 (0.72 1.11)	1.17 (0.62, 2.20)
Efavirenz b 600 mg QD (15 doses) 500/100 mg BID 24 ↔ 1.09 (0.99, 1.19) 1.04 (0.97, 1.12) 1.02 (0.92, 1.12) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 1.04) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 1.04) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 1.04) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 1.04) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 1.04) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 1.04) (15 doses) 1.12 (0.98, 1.28) 1.00 (0.93, 1.09) 0.94 (0.84, 0.56) 1.02 (0.48, 0.56) 1.02 (0.48, 0.56) 1.02 (0.48, 0.56) 1.02 (0.48, 0.56) 1.02 (0.48, 0.56) 1.02 (0.48, 0.42, 0.57) 0.57 (0.54, 0.60) 1.02 (0.94, 1.01) 0.99 (0.97, 1.02) 0.98 (0.94, 1.02) 1.00 mg QD (0.94, 0.94, 0.94, 0.94, 0.94, 0.94, 0.94, 0.94, 0.95) 0.89 (0.85, 0.92) 1.02 (0.94, 0				3	$\leftrightarrow$	0.80 (0.03, 1.02)	0.90 (0.72, 1.11)	1.17 (0.02, 2.20)
Ethinyl estradiol (15 doses) 750/200 mg BID (15 doses) 22	БС . В			24	$\leftrightarrow$	1.09 (0.99, 1.19)	1.04 (0.97, 1.12)	1.02 (0.92, 1.12)
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Efavirenz		750/200 mg BID			, , , ,		
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Ethinyl estradiol	0.035 mg	500/100 mg BID	21	$\overline{}$	0.52 (0.47, 0.57)	0.52 (0.48, 0.56)	•
then (2 or 14 doses) 19 $\leftrightarrow$ 0.94 (0.91, 0.98) 0.92 (0.88, 0.95) 0.89 (0.85, 0.92) (0.86, 0.92) 0.89 (0.85, 0.92) 0.89 (		(1 dose)		13	<b>1</b>	0.48 (0.42, 0.57)	0.57 (0.54, 0.60)	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Fluconazole	200 mg (Day 1)	500/200 mg BID		$\leftrightarrow$			
Lopinavir/RTV $= \begin{bmatrix} 400/100 \text{ mg BID} \\ (27 \text{ doses}) \end{bmatrix} = \begin{bmatrix} 500/200 \text{ mg BID} \\ (28 \text{ doses}) \end{bmatrix} = \begin{bmatrix} 21 \\ 69 \\ 400/100 \text{ mg BID} \end{bmatrix} = \begin{bmatrix} 0.53 \\ (0.40, 0.69) \end{bmatrix} = \begin{bmatrix} 0.45 \\ (0.32, 0.63) \end{bmatrix} = \begin{bmatrix} 0.30 \\ (0.17, 0.51) \end{bmatrix} = \begin{bmatrix} 0.48 \\ (0.40, 0.58) \end{bmatrix} = \begin{bmatrix} 0.48 \\ (0.40, 0.61) \end{bmatrix} = \begin{bmatrix} 0.48 \\ (0$		100 mg QD	(2 or 14 doses)	19	$\leftrightarrow$	0.94 (0.91, 0.98)	0.92 (0.88, 0.95)	0.89 (0.85, 0.92)
Loperamide 16 mg 750/200 mg BID 24 \$\frac{1}{2}\$ 0.39 (0.31, 0.48) 0.49 (0.40, 0.61) 0.48 (0.40, 0.58) 0.49 (0.40, 0.61)	8		500/200 ma DID	21	1	a a	d	d
Loperamide 16 mg 750/200 mg BID 24 ↓ 0.39 (0.31, 0.48) 0.49 (0.40, 0.61) (1 dose) (21 doses)	Lopinavir/RTV		•		Ť	0.53 (0.40, 0.69)	0.45 (0.32, 0.63)	0.30 (0.17, 0.51)
(1 dose) (21 doses)		(27 40303)	(20 00303)	- 07	*	•	-	0.48 (0.40, 0.58)
N-Demethyl-Loperamide 24 \ 0.21 (0.17, 0.25) 0.23 (0.19, 0.27)	Loperamide	•	•	24	1	0.39 (0.31, 0.48)	0.49 (0.40, 0.61)	-
	N-Demethyl-Lop	eramide		24	1	0.21 (0.17, 0.25)	0.23 (0.19, 0.27)	

<sup>&</sup>quot;HIV+ patients (TPV/ritonavir 250 mg/200 mg, 750 mg/200 mg and 1250 mg/100 mg) and healthy volunteers (TPV/ritonavir 500 mg/100 mg and 750 mg/200 mg)

"Normalized sum of parent drug (rifabutin) and active metabolite (25-O-desacetyl-rifabutin)

dIntensive PK analysis
"Drug levels obtained at 8-16 hrs post-dose

Drug Interactions: Pharmacokinetic Parameters for Co-administered Table 5 Drug in the Presence of tipranavir/ritonavir (continued)

Co- administered	Co- administered	TPV/ritonavir		DV/ -		ence Interval) of Co-ad arameters with/withou No Effect = 1.00	
Drug	Drug Dose (Schedule)	Drug Dose (Schedule)	D	PK -	C <sub>max</sub>	AUC	C <sub>min</sub>
Lamivudine a	150 mg BID (43 doses)	250/200 mg BID 750/100 mg BID	64	$\leftrightarrow$	0.96 (0.89, 1.03)	0.95 (0.89, 1.02)	-
	,	1250/100 mg BID (42 doses)	46	$\leftrightarrow$	0.86 (0.78, 0.94)	0.96 (0.90, 1.03)	-
			35	$\leftrightarrow$	0.71 (0.62, 0.81)	0.82 (0.66, 1.00)	-
Navionia a	200 mg BID	250/200 mg BID	26	$\leftrightarrow$	0.97 (0.90, 1.04)	0.97 (0.91, 1.04)	0.96 (0.87, 1.05)
Nevirapine	(43 doses)	750/100 mg BID	22	$\leftrightarrow$	0.86 (0.76, 0.97)	0.89 (0.78, 1.01)	0.93 (0.80, 1.08)
	, ,	1250/100 mg BID (42 doses)	17	$\leftrightarrow$	0.71 (0.62, 0.82)	0.76 (0.63, 0.91)	0.77 (0.64, 0.92)
Norethindrone	1.0 mg	500/100 mg BID	21	$\leftrightarrow$	1.03 (0.94, 1.13)	1.14 (1.06,1.22)	-
	(1 dose)	750/200 mg BID (21 doses)	13	$\leftrightarrow$	1.08 (0.97, 1.20)	1.27 (1.13,1.43)	-
Rifabutin	150 mg (1 dose)	500/200 mg BID (15 doses)	20	<b>↑</b>	1.70 (1.49, 1.94)	2.90 (2.59, 3.26)	2.14 (1.90, 2.41)
25-O-desacetyl-r	ifabutin	•	20	1	3.20 (2.78, 3.68)	20.71 (17.66, 24.28)	7.83 (6.70, 9.14)
Rifabutin + 25-C	-desacetyl-	•	20	Ť	1.86 (1.63, 2.12)	4.33 (3.86, 4.86)	2.76 (2.44, 3.12)
а	600/100 mg	500/200 mg BID	20	$\overline{}$	$0.30 (0.23, 0.40)^{d}$	0.24 (0.10, 0.22)	0.19(0.13.0.36)
Saquinavir/RTV	BID	(28 doses)	68	Ĭ	•	0.24 (0.19, 0.32)	0.18(0.13,0.26) e
	(27 doses)	` ,		•		-	0.20(0.16,0.25)
Stavudine <sup>a</sup>	40 mg BID,	250/200 mg BID	26	$\leftrightarrow$	0.90 (0.81, 1.02)	1.00 (0.91, 1.11)	-
Stavudine	≥60 kg	750/100 mg BID	22	$\leftrightarrow$	0.76 (0.66, 0.89)	0.84 (0.74, 0.96)	-
	30 mg B1D, <60 kg (43 doses)	1250/100 mg BID (42 doses)	19	$\leftrightarrow$	0.74 (0.69, 0.80)	0.93 (0.83, 1.05)	-
Tenofovir	300 mg	500/100 mg BID	22	Ţ	0.77 (0.68, 0.87)	0.98 (0.91, 1.05)	1.07 (0.98, 1.17)
Tenotovii	(1 dose)	750/200 mg BID (23 doses)	20	Ť	. 0.62 (0.54, 0.71)	1.02 (0.94, 1.10)	1.14 (1.01, 1.27)
Zidovudine	300 mg BID	250/200 mg BID	48	$\overline{}$	0.54 (0.47, 0.62)	0.58 (0.51, 0.66)	-
Lidovudine	300 mg BID	750/100 mg BID	31	Ţ	0.51 (0.44, 0.60)	0.64 (0.55, 0.75)	-
	300 mg BID	1250/100 mg BID	23	Ĭ	0.49 (0.40, 0.59)	0.69 (0.49, 0.97)	-
	(43 doses)	(42 doses)					
	300 mg	500/100 mg BID	29	<b>—</b>	0.39 (0.33, 0.45)	0.57 (0.52, 0.63)	0.89 (0.81, 0.99)
	(1 dose)	750/200 mg BID (23 doses)	25	ţ.	0.44 (0.36, 0.54)	0.67 (0.62, 0.73)	1.25 (1.08, 1.44)
Zidovudine gluci	uronide	500/100 mg BID	29	1	0.82 (0.74, 0.90)	1.02 (0.97, 1.06)	1.52 (1.34, 1.71)
_		750/200 mg BID (23 doses)	25	1	0.82 (0.73, 0.92)	1.09 (1.05, 1.14)	1.94 (1.62, 2.31)

<sup>\*</sup>HIV+ patients

bHIV+ patients (TPV/ritonavir 250 mg/200 mg, 750 mg/200 mg and 1250 mg/100 mg) and healthy volunteers (TPV/ritonavir 500 mg/100 mg and 750 mg/200 mg)

shormalized sum of parent drug (rifabutin) and active metabolite (25-O-desacetyl-rifabutin)

dIntensive PK analysis

Drug levels obtained at 8-16 hrs post-dose

#### INDICATIONS AND USAGE

APTIVUS® (tipranavir), co-administered with 200 mg of ritonavir, is indicated for combination antiretroviral treatment of HIV-1 infected adult patients with evidence of viral replication, who are highly treatment-experienced or have HIV-1 strains resistant to multiple protease inhibitors.

This indication is based on analyses of plasma HIV-1 RNA levels in two controlled studies of APTIVUS/ritonavir of 24 weeks duration. Both studies were conducted in clinically advanced, 3-class antiretroviral (NRTI, NNRTI, PI) treatment-experienced adults with evidence of HIV-1 replication despite ongoing antiretroviral therapy.

The following points should be considered when initiating therapy with APTIVUS/ritonavir:

- The use of other active agents with APTIVUS/ritonavir is associated with a greater likelihood of treatment response (see CLINICAL PHARMACOLOGY, Microbiology and INDICATIONS AND USAGE, Description of Clinical Studies).
- Genotypic or phenotypic testing and/or treatment history should guide the use of APTIVUS/ritonavir (see CLINICAL PHARMACOLOGY, Microbiology). The number of baseline primary protease inhibitor mutations affects the virologic response to APTIVUS/ritonavir (see CLINICAL PHARMACOLOGY, Microbiology).
- Liver function tests should be performed at initiation of therapy with APTIVUS/ritonavir and monitored frequently throughout the duration of treatment (see WARNINGS).
- Use caution when prescribing APTIVUS/ritonavir to patients with elevated transaminases, hepatitis B or C co-infection or other underlying hepatic impairment (see WARNINGS).
- The extensive drug-drug interaction potential of APTIVUS/ritonavir when co-administered with multiple classes of drugs must be considered prior to and during APTIVUS/ritonavir use (see CLINICAL PHARMACOLOGY and CONTRAINDICATIONS).
- The risk-benefit of APTIVUS/ritonavir has not been established in treatment-naïve adult patients or pediatric patients.

There are no study results demonstrating the effect of APTIVUS/ritonavir on clinical progression of HIV-1.

## **Description of Clinical Studies**

The following clinical data is derived from analyses of 24-week data from ongoing studies measuring effects on plasma HIV-1 RNA levels and CD4+ cell counts. At present there are no results from controlled studies evaluating the effect of APTIVUS/ritonavir on clinical progression of HIV.

## Treatment-Experienced Patients

Studies 1182.12 and 1182.48: APTIVUS/ritonavir 500/200 mg BID + optimized background regimen (OBR) vs. Comparator Protease Inhibitor/ritonavir BID + OBR

Studies 1182.12 and 1182.48 are ongoing, randomized, controlled, open-label, multicenter studies in HIV-positive, triple antiretroviral class experienced patients. All patients were required to have previously received at least two protease inhibitor-based antiretroviral regimens and were failing a protease inhibitor-based regimen at the time of study entry with baseline HIV-1 RNA at least 1000 copies/mL and any CD4+ cell count. At least one primary protease gene mutation from among 30N, 46I, 46L, 48V, 50V, 82A, 82F, 82L, 82T, 84V or 90M had to be present at baseline, with not more than two mutations at codons 33, 82, 84 or 90.

These studies evaluated treatment response at 24 weeks in a total of 1159 patients receiving either APTIVUS co-administered with 200 mg of ritonavir plus OBR versus a control group receiving a ritonavir-boosted protease inhibitor (lopinavir, amprenavir, saquinavir or indinavir) plus OBR. Prior to randomization, OBR was individually defined for each patient based on genotypic resistance testing and patient history. The investigator had to declare OBR, comparator protease inhibitor, and use of enfuvirtide prior to randomization. Randomization was stratified by choice of comparator protease inhibitor and use of enfuvirtide.

After Week 8, patients in the control group who met the protocol defined criteria of initial lack of virologic response had the option of discontinuing treatment and switching over to APTIVUS/ritonavir in a separate roll-over study.

Demographics and baseline characteristics were balanced between the APTIVUS/ritonavir arm and control arm. In both studies combined, the 1159 patients had a median age of 43 years (range 17-80), were 88% male, 73% white, 14% black and 1% Asian. The median baseline plasma HIV-1 RNA was 4.82 (range 2 to 6.8)  $\log_{10}$  copies/mL and median baseline CD4+ cell count was 155 (range 1 to 1893) cells/mm<sup>3</sup>. Forty percent (40%) of the patients had baseline HIV-1 RNA of  $\geq$  100,000 copies/mL, 61% had a baseline CD4+ cell count < 200 cells/mm<sup>3</sup>, and 57% had experienced an AIDS defining Class C event at baseline.

Patients had prior exposure to a median of 6 NRTIs, 1 NNRTI, and 4 PIs. A total of 12% of patients had previously used enfuvirtide. In baseline patient samples (n=454), 97% of the isolates were resistant to at least one protease inhibitor, 95% of the isolates were resistant to at least one NRTI, and > 75% of the isolates were resistant to at least one NNRTI.

The individually pre-selected protease inhibitor based on genotypic testing and the patient's medical history was lopinavir in 50%, amprenavir in 26%, saquinavir in 20% and indinavir in 4% of patients. A total of 86% were possibly resistant or resistant to the pre-selected comparator protease inhibitors. Approximately 25% of patients used enfuvirtide during study. There were differences between Studies 1182.12 and 1182.48 in the use of the protease inhibitors and in the use of enfuvirtide.

Treatment response and efficacy outcomes of randomized treatment through Week 24 of Studies 1182.12 and 1182.48 are shown in Table 6.

Table 6 Outcomes of Randomized Treatment Through Week 24 (Pooled Studies 1182.12 and 1182.48)

Outcome	Tipranavir/ritonavir (500/200 mg BID) + OBR (N = 582)	Comparator Protease Inhibitor*/ritonavir + OBR (N = 577)
Virological Responders <sup>a</sup> (confirmed at least 1 log <sub>10</sub> HIV-1	40%	18%
RNA below baseline)		
Virological failures Initial lack of virologic	54%	79%
response by Week 8 <sup>b</sup>	35%	59%
Rebound	12%	11%
Never suppressed	7%	8%
Death <sup>c</sup> or discontinued due to		
adverse events	1%	1%
Discontinued due to other reasons	5%	2%

<sup>\*</sup>Comparator protease inhibitors were lopinavir, amprenavir, saquinavir or indinavir and 86% of patients were possibly resistant or resistant to the chosen protease inhibitors.

Through 24 weeks of treatment, the proportion of patients in the APTIVUS/ritonavir arm compared to the comparator PI/ritonavir arm with HIV-1 RNA < 400 copies/mL was 34% and 16% respectively, and with HIV-1 RNA < 50 copies/mL was 23% and 9% respectively. Among all randomized and treated patients, the median change from baseline in HIV-1 RNA at the last measurement up to Week 24 was -0.80 log<sub>10</sub> copies/mL in patients receiving APTIVUS/ritonavir versus -0.25 log<sub>10</sub> copies/mL in the comparator PI/ritonavir arm.

Among all randomized and treated patients, the median change from baseline in CD4+ cell count at the last measurement up to Week 24 was +34 cells/mm<sup>3</sup> in patients receiving tipranavir/ritonavir (N = 582) versus +4 cells/mm<sup>3</sup> in the comparator PI/ritonavir (N = 577) arm.

Patients in the APTIVUS/ritonavir arm achieved a significantly better virologic outcome when APTIVUS/ritonavir was combined with enfuvirtide (see CLINICAL PHARMACOLOGY, Microbiology).

#### CONTRAINDICATIONS

APTIVUS (tipranavir) is contraindicated in patients with known hypersensitivity to any of the ingredients of the product.

APTIVUS is contraindicated in patients with moderate and severe (Child-Pugh Class B and C, respectively) hepatic insufficiency (see WARNINGS).

<sup>&</sup>lt;sup>a</sup>Patients achieved and maintained a confirmed ≥ 1 log<sub>10</sub> HIV-1 RNA drop from baseline through Week 24 without prior evidence of treatment failure.

<sup>&</sup>lt;sup>b</sup>Patients did not achieve a 0.5 log HIV-1 RNA drop from baseline and did not have viral load < 100,000 copies/mL by Week 8.

<sup>&</sup>lt;sup>c</sup>Patients who died while being virologically suppressed.

dIncludes patients who were lost to-follow-up, withdrawn consent, non-adherent, protocol violations, added/changed background antiretroviral drugs for reasons other than tolerability or toxicity, or discontinued while suppressed.

Co-administration of APTIVUS with 200 mg of ritonavir with drugs that are highly dependent on CYP 3A for clearance and for which elevated plasma concentrations are associated with serious and/or life-threatening events is contraindicated. These drugs are listed in Table 7 below. For information regarding clinical recommendations see PRECAUTIONS, Drug Interactions, Tables 8 and 9.

Table 7 Drugs that are Contraindicated with Tipranavir, Co-Administered with 200 mg of Ritonavir

Drug Class	Drugs within Class that are Contraindicated with APTIVUS Co-administered with 200 mg of ritonavir
Antiarrhythmics	Amiodarone, bepridil, flecainide, propafenone, quinidine
Intihistamines	Astemizole, terfenadine
Ergot derivatives	Dihydroergotamine, ergonovine, ergotamine, methylergonovine
I motility agent	Cisapride
Neuroleptic	Pimozide
Sedatives/hypnotics	Midazolam, triazolam

Due to the need for co-administration of APTIVUS with 200 mg of ritonavir, please refer to ritonavir prescribing information for a description of ritonavir contraindications.

#### WARNINGS

ALERT: Find out about medicines that should NOT be taken with APTIVUS. This statement is included on the product's bottle label.

APTIVUS (tipranavir) must be co-administered with 200 mg of ritonavir to exert its therapeutic effect (see **DOSAGE AND ADMINISTRATION**). Failure to correctly co-administer APTIVUS with ritonavir will result in reduced plasma levels of tipranavir that will be insufficient to achieve the desired antiviral effect and will alter some drug interactions (effect of tipranavir and ritonavir on other drugs).

Please refer to ritonavir prescribing information for additional information on precautionary measures.

# **Hepatic Impairment and Toxicity**

APTIVUS co-administered with 200 mg of ritonavir, has been associated with reports of clinical hepatitis and hepatic decompensation, including some fatalities. These have generally occurred in patients with advanced HIV disease taking multiple concomitant medications. A causal relationship to APTIVUS/ritonavir could not be established. All patients should be followed closely with clinical and laboratory monitoring, especially those with chronic hepatitis B or C co-infection, as these patients have an increased risk of hepatotoxicity. Liver function tests should be performed prior to initiating therapy with APTIVUS/ritonavir, and frequently throughout the duration of treatment.

Patients with chronic hepatitis B or hepatitis C co-infection or elevations in transaminases are at approximately 2.5-fold risk for developing further transaminase elevations or hepatic decompensation. Additionally, Grade 3 and 4 increases in hepatic transaminases were observed in 6% of healthy volunteers in Phase 1 studies and 6% of subjects receiving APTIVUS/ritonavir in Phase 3 studies.

Tipranavir is principally metabolized by the liver. Therefore caution should be exercised when administering APTIVUS/ritonavir to patients with hepatic impairment because tipranavir concentrations may be increased. APTIVUS/ritonavir is contraindicated in patients with moderate to severe (Child-Pugh Class B and Child-Pugh Class C) hepatic insufficiency.

Physicians and patients should be vigilant for the appearance of signs or symptoms of hepatitis, such as fatigue, malaise, anorexia, nausea, jaundice, bilirubinuria, acholic stools, liver tenderness or hepatomegaly. Patients with signs or symptoms of clinical hepatitis should discontinue APTIVUS/ritonavir treatment and seek medical evaluation.

For information on the multi-dose pharmacokinetics of tipranavir in hepatically impaired patients (see CLINICAL PHARMACOLOGY, Pharmacokinetics in Special Populations, *Hepatic Impairment*).

# Diabetes Mellitus/Hyperglycemia

New onset diabetes mellitus, exacerbation of pre-existing diabetes mellitus and hyperglycemia have been reported during post-marketing surveillance in HIV-1 infected patients receiving protease inhibitor therapy. Some patients required either initiation or dose adjustments of insulin or oral hypoglycemic agents for treatment of these events. In some cases, diabetic ketoacidosis has occurred. In those patients who discontinued protease inhibitor therapy, hyperglycemia persisted in some cases. Because these events have been reported voluntarily during clinical practice, estimates of frequency cannot be made and a causal relationship between protease inhibitor therapy and these events has not been established.

#### **PRECAUTIONS**

#### Sulfa Allergy

APTIVUS (tipranavir) should be used with caution in patients with a known sulfonamide allergy. Tipranavir contains a sulfonamide moiety. The potential for cross-sensitivity between drugs in the sulfonamide class and tipranavir is unknown.

#### Rash

Mild to moderate rashes including urticarial rash, maculopapular rash, and possible photosensitivity have been reported in subjects receiving APTIVUS/ritonavir. In Phase 2 and 3 trials rash was observed in 14% of females and in 8-10% of males receiving APTIVUS/ritonavir. Additionally, in one drug interaction trial in healthy female volunteers administered a single dose of ethinyl estradiol followed by APTIVUS/ritonavir, 33% of subjects developed a rash. Rash accompanied by joint pain or stiffness, throat tightness, or generalized pruritus has been reported in both men and women receiving APTIVUS/ritonavir (see PRECAUTIONS, Drug Interactions and ADVERSE REACTIONS).

#### Patients with Hemophilia

There have been reports of increased bleeding, including spontaneous skin hematomas and hemarthrosis in patients with hemophilia type A and B treated with protease inhibitors. In some patients additional Factor VIII was given. In more than half of the reported cases, treatment with

protease inhibitors was continued or reintroduced if treatment had been discontinued. A causal relationship between protease inhibitors and these events has not been established.

# **Lipid Elevations**

Treatment with APTIVUS co-administered with 200 mg of ritonavir has resulted in large increases in the concentration of total cholesterol and triglycerides (see ADVERSE REACTIONS, Table 11). Triglyceride and cholesterol testing should be performed prior to initiating APTIVUS/ritonavir therapy and at periodic intervals during therapy. Lipid disorders should be managed as clinically appropriate (see PRECAUTIONS, Drug Interactions, Table 9: Established and Other Potentially Significant Drug Interactions for additional information on potential drug interactions with APTIVUS/ritonavir and HMG-CoA reductase inhibitors).

## **Fat Redistribution**

Redistribution/accumulation of body fat including central obesity, dorsocervical fat enlargement (buffalo hump), peripheral wasting, facial wasting, breast enlargement, and "cushingoid appearance" have been observed in patients receiving antiretroviral therapy. The mechanism and long-term consequences of these events are currently unknown. A causal relationship has not been established.

#### **Immune Reconstitution Syndrome**

Immune reconstitution syndrome has been reported in patients treated with combination antiretroviral therapy, including tipranavir. During the initial phase of combination antiretroviral treatment, patients whose immune system responds may develop an inflammatory response to indolent or residual opportunistic infections (such as *Mycobacterium avium* infection, cytomegalovirus, *Pneumocystis jeroveci* pneumonia, tuberculosis, or reactivation of herpes simplex and herpes zoster), which may necessitate further evaluation and treatment.

#### **Information for Patients**

Patients should be informed that APTIVUS co-administered with 200 mg of ritonavir, has been associated with severe liver disease, including some deaths. Patients with signs or symptoms of clinical hepatitis should discontinue APTIVUS/ritonavir treatment and seek medical evaluation. Symptoms of hepatitis include fatigue, malaise, anorexia, nausea, jaundice, bilirubinuria, acholic stools, liver tenderness or hepatomegaly. Extra vigilance is needed for patients with chronic hepatitis B or C co-infection, as these patients have an increased risk of hepatotoxicity.

Liver function tests should be performed prior to initiating therapy with tipranavir and 200 mg of ritonavir, and frequently throughout the duration of treatment. Patients with chronic hepatitis B or C co-infection or elevations in liver enzymes prior to treatment are at increased risk (approximately 2.5-fold) for developing further liver enzyme elevations or severe liver disease. Caution should be exercised when administering APTIVUS/ritonavir to patients with liver enzyme abnormalities or history of chronic liver disease. Increased liver function testing is warranted in these patients. APTIVUS should not be given to patients with moderate to severe liver disease.

Mild to moderate rash has been reported in HIV-infected men and women receiving APTIVUS/ritonavir.

Women receiving estrogen-based hormonal contraceptives should be instructed that additional or alternative contraceptive measures should be used during therapy with APTIVUS/ritonavir. There may be an increased risk of rash when APTIVUS is given with hormonal contraceptives.

Patients should be informed that redistribution or accumulation of body fat may occur in patients receiving antiretroviral therapy and that the cause and long-term health effects of these conditions are not known at this time.

Patients should be informed that APTIVUS must be co-administered with 200 mg ritonavir to ensure its therapeutic effect. Failure to correctly co-administer APTIVUS with ritonavir will result in reduced plasma levels of tipranavir that may be insufficient to achieve the desired antiviral effect.

Patients should be told that sustained decreases in plasma HIV-1 RNA have been associated with a reduced risk of progression to AIDS and death. Patients should remain under the care of a physician while using APTIVUS. Patients should be advised to take APTIVUS and other concomitant antiretroviral therapy every day as prescribed. APTIVUS, co-administered with ritonavir, must be given in combination with other antiretroviral drugs. Patients should not alter the dose or discontinue therapy without consulting with their doctor. If a dose of APTIVUS is missed, patients should take the dose as soon as possible and then return to their normal schedule. However, if a dose is skipped the patient should not double the next dose.

Patients should be informed that APTIVUS is not a cure for HIV-1 infection and that they may continue to develop opportunistic infections and other complications associated with HIV disease. The long-term effects of APTIVUS are unknown at this time. Patients should be told that there are currently no data demonstrating that therapy with APTIVUS can reduce the risk of transmitting HIV to others through sexual contact.

APTIVUS may interact with some drugs; therefore, patients should be advised to report to their health care provider the use of any other prescription, non-prescription medication or herbal products, particularly St. John's wort.

APTIVUS should be taken with food to enhance absorption.

The Patient Package Insert provides written information for the patients, and should be dispensed with each new prescription and refill.

## **Drug Interactions**

Tipranavir administered with ritonavir can alter plasma exposure of other drugs and other drugs can alter plasma exposure of tipranavir and ritonavir.

Tipranavir co-administered with 200 mg of ritonavir at the recommended dosage is a net inhibitor of CYP 3A and may increase plasma concentrations of agents that are primarily metabolized by CYP 3A. Thus, co-administration of tipranavir/ritonavir with drugs highly dependent on CYP 3A for clearance and for which elevated plasma concentrations are associated with serious and/or life-threatening events should be contraindicated. Co-administration with other CYP 3A substrates may require a dose adjustment or additional monitoring (see CONTRAINDICATIONS and PRECAUTIONS).

The mechanisms of the potential interactions are described in the CLINICAL PHARMACOLOGY, Drug Interactions section.

Drugs that are contraindicated or not recommended for co-administration with APTIVUS are included in Table 8 below. These recommendations are based on either drug interaction studies or they are predicted interactions due to the expected magnitude of interaction and potential for serious events or loss of efficacy.

Table 8 Drugs that Should Not be Co-administered with APTIVUS Co-administered with 200 mg of Ritonavir

Drug Class/Drug Name	Clinical Comment
Antiarrhythmics Amiodarone, bepridil, flecainide, propafenone, quinidine	CONTRAINDICATED due to potential for serious and/or life- threatening reactions such as cardiac arrhythmias secondary to increases in plasma concentrations of antiarrhythmics.
Antihistamines Astemizole, terfenadine	CONTRAINDICATED due to potential for serious and/or life- threatening reactions such as cardiac arrhythmias.
Antimycobacterials Rifampin	May lead to loss of virologic response and possible resistance to tipranavir or to the class of protease inhibitors.
Ergot derivatives Dihydroergotamine, ergonovine, ergotamine, methylergonovine	CONTRAINDICATED due to potential for serious and/or life- threatening reactions such as acute ergot toxicity characterized by peripheral vasospasm and ischemia of the extremities and other tissues.
GI motility agents Cisapride	CONTRAINDICATED due to potential for serious and/or life- threatening reactions such as cardiac arrhythmias.
Herbal products St. John's wort	May lead to loss of virologic response and possible resistance to tipranavir or to the class of protease inhibitors.
HMG CoA reductase inhibitors Lovastatin, simvastatin	Potential for serious reactions such as risk of myopathy including rhabdomyolysis.
Neuroleptics Pimozide	CONTRAINDICATED due to potential for serious and/or life- threatening reactions such as cardiac arrhythmias.
Sedatives/hypnotics Midazolam, triazolam	CONTRAINDICATED due to potential for serious and/or life threatening reactions such as prolonged or increased sedation or respiratory depression.

Clinically significant drug-drug interactions of APTIVUS co-administered with 200 mg of ritonavir are summarized in the Table 9 below.

Table 9 Established and Other Potentially Significant Drug Interactions:
Alterations in Dose or Regimen May be Recommended Based on Drug
Interaction Studies or Predicted Interaction (continued)

Concomitant Drug Class: Drug name	Effect on Concentration of Tipranavir or Concomitant Drug	Clinical Comment
	Other Agents for Opportunistic Infecti	ons
Antimycobacterials:		
Clarithromycin	↑ Tipranavir, ↑ Clarithromycin, ↓ 14-hydroxy-clarithromycin metabolite	No dose adjustment of tipranavir or clarithromycin for patients with normal renal function is necessary.
	•	For patients with renal impairment the following dosage adjustments should be considered:
		<ul> <li>For patients with CL<sub>CR</sub> 30 to 60 mL/min the dose of clarithromycin should be reduced by 50%.</li> </ul>
		For patients with CL <sub>CR</sub> < 30 mL/min the dose of clarithromycin should be decreased by 75%.
Rifabutin	Tipranavir not changed, ↑Rifabutin ↑ Desacetyl-rifabutin	Single dose study. Dosage reductions of rifabutin by 75% are recommended (e.g. 150 mg every other day). Increased monitoring for adverse events in patients receiving the combination is warranted. Further dosage reduction may be necessary.
	Other Agents Commonly used	
Calcium Channel Blockers:	Combination with TPV/ritonavir	Caution is warranted and clinical
Diltiazem	not studied. Cannot predict effect of TPV/ritonavir on calcium	monitoring of patients is recommended.
Felodipine '	channel blockers that are dual	
Nicardipine	substrates of CYP 3A and P-gp due	
Nisoldipine	to conflicting effect of	
Verapamil	TPV/ritonavir on CYP 3A and P-	
	gp.  ↑ Diltiazem  ↑ Felodipine (CYP 3A	
	substrate but not P-gp substrate)  † Nicardipine  † Nisoldipine (CYP 3A substrate	
	but not clear whether it is a P-gp substrate)  1 Verapamil	
Despiramine	Combination with TPV/ritonavir not studied  ↑ Despiramine	Dosage reduction and concentration monitoring of despiramine is recommended.

Table 9 Established and Other Potentially Significant Drug Interactions:
Alterations in Dose or Regimen May be Recommended Based on Drug
Interaction Studies or Predicted Interaction (continued)

Disulfiram/Metronidazole	Combination with TPV/ritonavir not studied	APTIVUS capsules contain alcohol that can produce disulfiram-like reactions when co-administered with disulfiram or other drugs which produce this reaction (e.g. metronidazole).
HMG-CoA reductase inhibitors:		Start with the lowest possible dose
Atorvastatin	↑ Tipranavir, ↑ Atorvastatin ↓ Hydroxy-atorvastatin metabolites	of atorvastatin with careful monitoring, or consider other HMG-CoA reductase inhibitors. Concomitant use of APTIVUS, coadministered with 200 mg of ritonavir, with lovastatin or simvastatin is not recommended.
Hypoglycemics:	Combination with TPV/ritonavir	Careful glucose monitoring is
	not studied.	warranted.
Glimepiride Glipizide Glyburide Pioglitazone Repaglinide Tolbutamide	☐ Glimepiride (CYP 2C9) ☐ Glipizide (CYP 2C9) ☐ Glyburide (CYP 2C9) ☐ Pioglitazone (CYP 2C8 and CYP 3A4) ☐ Repaglinide (CYP 2C8 and CYP 3A4) ☐ Tolbutamide (CYP 2C9)  The effect of TPV/ritonavir on CYP 2C8 and CYP 2C9 substrates is not known.  Combination with TPV/ritonavir	More frequent concentration
Cyclosporine Sirolimus Tacrolimus	not studied. Cannot predict effect of TPV/ritonavir on immunosuppressants due to conflicting effect of TPV/ritonavir on CYP 3A and P-gp.  † Cyclosporine † Sirolimus † Tacrolimus	monitoring of these medicinal products is recommended until blood levels have been stabilized.
Narcotic analgesics:		
Meperidine	Combinations with TPV/ritonavir not studied  ↓ Meperidine, ↑ Normeperidine	Dosage increase and long-term use of meperidine are not recommended due to increased concentrations of the metabolite normeperidine which has both analgesic activity and CNS stimulant activity (e.g. seizures).
Methadone	↓ Methadone by 50% .	Dosage of methadone may need to be increased when co-administered with tipranavir and 200 mg of ritonavir.

Table 9 Established and Other Potentially Significant Drug Interactions:
Alterations in Dose or Regimen May be Recommended Based on Drug
Interaction Studies or Predicted Interaction (continued)

Oral contraceptives/Estrogens:		Alternative methods of nonhormonal contraception should be used when
Ethinyl estradiol	↓ Ethinyl estradiol concentrations by 50%	estrogen based oral contraceptives are co-administered with tipranavir and 200 mg of ritonavir. Patients using estrogens as hormone replacement therapy should be clinically monitored for signs of estrogen deficiency.  Women using estrogens may have an increased risk of non serious rash.
PDE5 inhibitors:	Combinations with TPV/ritonavir not studied.	Concomitant use of PDE5 inhibitors with tipranavir and ritonavir should
Sildenafil	↑ Sildenafil	be used with caution and in no case
Tadalafil	↑ Tadalafil	should the starting dose of:
Vardenafil	↑ Vardenafil	<ul> <li>sildenafil exceed 25 mg within 48 hours</li> <li>tadalafil exceed 10 mg every 72 hours</li> <li>vardenafil exceed 2.5 mg every 72 hours</li> </ul>
Selective Serotonin-Reuptake Inhibitors:	Combination with TPV/ritonavir not studied	Antidepressants have a wide therapeutic index, but doses may need to be adjusted upon initiation
Fluoxetine	↑ Fluoxetine	of APTIVUS/ritonavir therapy.
Paroxetine	↑ Paroxetine	
Sertraline	↑ Sertraline	
Warfarin	Combination with TPV/ritonavir not studied. Cannot predict the effect of TPV/ritonavir on S-Warfarin due to conflicting effect of TPV and RTV on CYP 2C9	Frequent INR (international normalized ratio) monitoring upon initiation of tipranavir/ritonavir therapy.

## Carcinogenesis, Mutagenesis, Impairment of Fertility

Long term animal carcinogenicity bioassays with tipranavir and tipranavir/ritonavir are currently in progress. However, tipranavir showed no evidence of mutagenicity or clastogenicity in a battery of five in vitro and in vivo tests including the Ames bacterial reverse mutation assay using S. typhimurium and E. coli, unscheduled DNA synthesis in rat hepatocytes, induction of gene mutation in Chinese hamster ovary cells, a chromosome aberration assay in human peripheral lymphocytes, and a micronucleus assay in mice.

Tipranavir had no effect on fertility or early embryonic development in rats at dose levels up to 1000 mg/Kg/day, equivalent to a  $C_{max}$  of 258  $\mu M$  in females. Based on  $C_{max}$  levels in these rats, as well as an exposure (AUC) of 1670  $\mu M$  h in pregnant rats from another study, this exposure was

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approximately equivalent to the anticipated exposure in humans at the recommended dose level of 500/200 mg tipranavir/ritonavir BID.

### Pregnancy

# Teratogenic Effects, Pregnancy Category C.

Investigation of fertility and early embryonic development with tipranavir disodium was performed in rats, teratogenicity studies were performed in rats and rabbits, and pre- and post-natal development were explored in rats.

No teratogenicity was detected in reproductive studies performed in pregnant rats and rabbits up to dose levels of 1000 mg/Kg/day and 150 mg/Kg/day tipranavir, respectively, at exposure levels approximately 1.1-fold and 0.1-fold human exposure. At 400 mg/Kg/day and above in rats, fetal toxicity (decreased sternebrae ossification and body weights) was observed, corresponding to an AUC of 1310  $\mu$ M·h or approximately 0.8-fold human exposure at the recommended dose. In rats and rabbits, fetal toxicity was not noted at 40 mg/Kg/day and 150 mg/Kg/day, respectively, corresponding accordingly to  $C_{max}/AUC_{0-24h}$  levels of 30.4  $\mu$ M/340  $\mu$ M·h and 8.4  $\mu$ M/120  $\mu$ M·h. These exposure levels (AUC) are approximately 0.2-fold and 0.1-fold the exposure in humans at the recommended dose.

In pre- and post-development studies in rats, tipranavir showed no adverse effects at 40 mg/Kg/day (~0.2-fold human exposure), but caused growth inhibition in pups and maternal toxicity at dose levels of 400 mg/Kg/day (~0.8-fold human exposure). No post-weaning functions were affected at any dose level.

There are no adequate and well-controlled studies in pregnant women for the treatment of HIV-1 infection. APTIVUS should be used during pregnancy only if the potential benefit justifies the potential risk to the fetus.

#### Antiretroviral Pregnancy Registry

To monitor maternal-fetal outcomes of pregnant women exposed to APTIVUS, an Antiretroviral Pregnancy Registry has been established. Physicians are encouraged to register patients by calling (800) 258-4263.

#### **Nursing Mothers**

The Centers for Disease Control and Prevention recommend that HIV-infected mothers not breastfeed their infants to avoid risking postnatal transmission of HIV. Because of both the potential for HIV transmission and any possible adverse effects of tipranavir, mothers should be instructed not to breastfeed if they are receiving APTIVUS.

#### **Pediatric Use**

Safety and effectiveness in pediatric patients have not been established.

#### Geriatric Use

Clinical studies of APTIVUS did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects. In general, caution should be exercised in the administration and monitoring of APTIVUS in elderly patients reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy.

## **ADVERSE REACTIONS**

APTIVUS (tipranavir), co-administered with 200 mg of ritonavir, has been studied in a total of 1854 HIV-positive adults as combination therapy in clinical studies. Of these, 1397 patients received the dose of 500/200 mg BID. Seven hundred sixty one (761) adults, including 385 in the 1182.12 and 1182.48 Phase 3 pivotal studies, have been treated for at least 24 weeks.

In 1182.12 and 1182.48 in the APTIVUS/ritonavir arm, the most frequent AEs were diarrhea, nausea, fatigue, headache and vomiting. Adverse events leading to discontinuation were reported by 7.8% of the tipranavir-treated patients and 4.9% of the comparator arm patients.

Due to the need for co-administration of APTIVUS with 200 mg of ritonavir, please refer to ritonavir prescribing information for ritonavir-associated adverse reactions.

The most frequent clinical treatment-emergent adverse events reported in Phase 3 clinical studies (1182.12 and 1182.48) in adults are summarized in Table 10 below. Events of moderate to severe intensity (Grades 2-4) reported in at least 2% of highly treatment-experienced subjects in either treatment group are included.

Table 10 Percentage of Patients with Treatment Emergent Adverse Events of at Least Moderate Intensity (Grades 2-4) in ≥ 2% of Patients in Either Treatment Group<sup>a</sup>

	Phase 3 Studies 1182.13	2 and 1182.48 (24-weeks)
	Tipranavir/ritonavir (500/200 mg BID) + OBR (n=746)	Comparator PI/ritonavir <sup>b</sup> + OBR (n=737)
Gastrointestinal Disorders		
Diarrhea	10.9%	9.4%
Nausea	6.7%	4.6%
Vomiting	3.4%	3.0%
Abdominal pain <sup>c</sup>	2.8%	3.7%
General Disorders		
Pyrexia	4.6%	4.3%
Fatigue	4.0%	3.9%
Asthenia	1.5%	2.3%
Infections and Infestations		
Bronchitis	2.9%	1.1%
Nervous System Disorders		
Headache	3.1%	3.1%
Psychiatric Disorders		
Depression	2.0%	3.0%
Insomnia	1.2%	2.6%
Respiratory, Thoracic and		
Mediastinal Disorders		
Cough	0.8%	2.2%
Skin and Subcutaneous Tissue		
Disorders		
Rash	2.0%	2.0%

<sup>&</sup>lt;sup>a</sup>Excludes laboratory abnormalities that were Adverse Events

Clinically meaningful adverse reactions in < 2% of adult patients (n=1397) treated with APTIVUS/ritonavir 500/200mg in Phase 2 and 3 trials listed below by body system:

Blood and Lymphatic System Disorders: anemia, neutropenia, thrombocytopenia

Gastrointestinal Disorders: abdominal distension, dyspepsia, flatulence, gastroesophageal reflux disease, pancreatitis

General Disorders: influenza like illness, malaise, pyrexia

Hepatobiliary Disorders: hepatitis, hepatic failure

Immune System Disorders: hypersensitivity

Infections and infestations: reactivation of herpes simplex and varicella zoster

Investigations: hepatic enzymes increased, liver function test abnormal, lipase increased, weight decreased

<sup>&</sup>lt;sup>b</sup>Comparator PI/RTV: lopinavir/ritonavir 400/100 mg BID, indinavir/ritonavir 800/100 mg BID, saquinavir/ritonavir 1000/100 mg BID, amprenavir/ritonavir 600/100 mg BID

<sup>&</sup>lt;sup>c</sup>Abdominal pain includes Preferred Terms "Abdominal pain" and "Abdominal pain upper"

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Metabolism and Nutrition Disorders: anorexia, decreased appetite, dehydration, diabetes mellitus, facial wasting, hyperamylasemia, hypercholesterolemia, hyperglycemia

Musculoskeletal and Connective Tissue Disorders: muscle cramp, myalgia

Nervous System Disorders: dizziness, neuropathy peripheral, somnolence

Psychiatric Disorders: insomnia, sleep disorder

Renal and Urinary Disorders: renal insufficiency

Respiratory, Thoracic and Mediastinal Disorders: dyspnea

Skin and Subcutaneous System Disorders: exanthem, lipoatrophy, lipodystrophy acquired, lipohypertrophy, pruritus

## **Laboratory Abnormalities**

Treatment-emergent clinical laboratory abnormalities reported at 24 weeks in Phase 3 clinical studies (1182.12 and 1182.48) in adults are summarized in Table 11 below.

Table 11 Treatment Emergent Laboratory Abnormalities Reported in ≥ 2% of Adult Patients

•		Studies 1182.12 a	nd 1182.48 (24-weeks)
	Limit	APTIVUS/ritonavir (500/200 mg BID) + OBR (n = 732)	Comparator PI/ritonavi + OBR* (n = 726)
Hematology			
· WBC count			
decrease			
Grade 3-4	< 2.0 x 10 <sup>3</sup> /μL	3.6%	5.4%
Chemistry			
Amylase			
Grade 3-4	> 2 x ULN	2.9%	4.8%
ALT			
Grade 2	> 2.5-5 x ULN	10.7%	5.4%
Grade 3	> 5-10 x ULN	3.1%	1.4%
Grade 4	> 10 x ULN	2.7%	0.4%
AST			
Grade 2	> 2.5-5 x ULN	6.0%	5.8%
Grade 3	> 5-10 x ULN	3.3%	1.0%
Grade 4	> 10 x ULN	0.7%	0.4%
ALT and/or AST			
Grade 2-4	> 2.5 x ULN	17.5%	9.9%
Cholesterol			
Grade 2	> 300 - 400  mg/dL	11.3%	4.3%
Grade 3	> 400 - 500  mg/dL	2.5%	0.3%
Grade 4	> 500 mg/dL	0.8%	0%
Triglycerides			
Grade 2	400 - 750  mg/dL	26.2%	14.7%
Grade 3	> 750 - 1200  mg/dL	12.8%	5.6%
Grade 4	> 1200 mg/dL	6.1%	3.4%

<sup>\*</sup>Comparator PI/RTV: lopinavir/ritonavir 400/100 mg BID, indinavir/ritonavir 800/100 mg BID, saquinavir/ritonavir 1000/100 mg BID, amprenavir/ritonavir 600/100 mg BID

In clinical trials extending up to 48 weeks, the proportion of patients who developed Grade 2-4 ALT and/or AST elevations increased to 24.4% with APTIVUS/ritonavir and to 12.8% with CPI/ritonavir.

## **OVERDOSAGE**

There is no known antidote for tipranavir overdose. Treatment of overdose should consist of general supportive measures, including monitoring of vital signs and observation of the patient's clinical status. If indicated, elimination of unabsorbed tipranavir should be achieved by emesis or gastric lavage. Administration of activated charcoal may also be used to aid in removal of unabsorbed drug. Since tipranavir is highly protein bound, dialysis is unlikely to be beneficial in significant removal of this medicine.

#### DOSAGE AND ADMINISTRATION

#### General

The recommended dose of APTIVUS (tipranavir) Capsules is 500 mg (two 250 mg capsules), coadministered with 200 mg of ritonavir, twice daily.

APTIVUS Capsules, co-administered with 200 mg of ritonavir should be taken with food. Bioavailability is increased with a high fat meal.

## **HOW SUPPLIED**

APTIVUS (tipranavir) Capsules 250 mg are pink, oblong soft gelatin capsules imprinted in black with "TPV 250". They are packaged in HDPE unit-of-use bottles with a child resistant closure and 120 capsules. (NDC 0597-0003-02)

APTIVUS capsules should be stored in a refrigerator 2°-8°C (36°-46°F) prior to opening the bottle. After opening the bottle, the capsules may be stored at 25°C (77°F); excursions permitted to 15°-30°C (59°-86°F) and must be used within 60 days.

Store in a safe place out of the reach of children.

Address medical inquiries to: <a href="http://us.boehringer-ingelheim.com">http://us.boehringer-ingelheim.com</a>, (800) 542-6257 or (800) 459-9906 TTY.

#### RX ONLY

Distributed by:

Boehringer Ingelheim Pharmaceuticals, Inc.

Ridgefield, CT 06877 USA

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APTIVUS Capsules are covered by U.S. Patents 5,852,195; 6,147,095; 6,169,181 and 6,231,887

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10003515/US/1

Revision Date: June 21, 2005

#### **Patient Information**

Aptivus®(ap'·ti·vəs) (tipranavir) Capsules, 250 mg



ALERT: Find out about medicines that should not be taken with Aptivus. Please also read the section "WHO SHOULD NOT TAKE APTIVUS".

Read the Patient Information that comes with APTIVUS before you start taking it and each time you get a refill. There may be new information. This leaflet does not take the place of talking with your doctor about your medical condition or treatment. You should stay under a doctor's care while taking APTIVUS.

## What is the most important information I should know about APTIVUS?

Patients taking APTIVUS, together with 200 mg NORVIR® (ritonavir), may develop severe liver disease that can cause death. If you develop any of the following symptoms of liver problems, you should stop taking APTIVUS/ritonavir treatment and call your doctor right away: tiredness, general ill feeling or "flu-like" symptoms, loss of appetite, nausea (feeling sick to your stomach), yellowing of your skin or whites of your eyes, dark (tea-colored) urine, pale stools (bowel movements), or pain, ache, or sensitivity on your right side below your ribs. If you have chronic Hepatitis B or C infection, your doctor should check your blood tests more often because you have an increased chance of developing liver problems.

## What is APTIVUS?

APTIVUS is a medicine called a "protease inhibitor" that is used to treat adults with Human Immunodeficiency Virus (HIV). APTIVUS blocks HIV protease, an enzyme which is needed for HIV to make more virus. When used with other anti-HIV medicines, APTIVUS may reduce the amount of HIV in your blood and increase the number of CD4+ cells. Reducing the amount of HIV in the blood may keep your immune system healthy, so it can help fight infection.

APTIVUS is always taken with NORVIR® (ritonavir) and at the same time as NORVIR When you take APTIVUS with NORVIR, you must always use at least 2 other anti-HIV medicines.

## Does APTIVUS cure HIV or AIDS?

APTIVUS does not cure HIV infection or AIDS. The long-term effects of APTIVUS are not known at this time. People taking APTIVUS may still get infections or other conditions common in people with HIV (opportunistic infections). It is very important that you stay under the care of your doctor during treatment with APTIVUS.

## Does APTIVUS lower the chance of passing HIV to other people?

APTIVUS does not reduce the chance of passing HIV to others through sexual contact, sharing needles, or being exposed to your blood. Continue to practice safer sex. Use a latex or polyurethane condom or other barrier method to lower the chance of sexual contact with any body fluids such as semen, vaginal secretions or blood. Never use or share dirty needles.

Ask your doctor if you have any questions about safer sex or how to prevent passing HIV to other people.

## Who should not take APTIVUS?

## Do not take APTIVUS if you:

- are allergic to tipranavir or any of the other ingredients in APTIVUS. See the end of this leaflet for a list of major ingredients.
- are allergic to ritonavir (NORVIR®)
- have moderate to severe liver problems
- take any of the following types of medicines because you could have serious side effects:
  - o Migraine headache medicines called "ergot alkaloids". If you take migraine headache medicines, ask you doctor or pharmacist if any of them are "ergot alkaloids".
  - o Halcion® (triazolam)
  - o Hismanal® (astemizole)
  - o Orap® (pimozide)
  - o Propulsid® (cisapride)
  - o Seldane® (terfenadine)
  - o Versed® (midazolam)
  - o Pacenone® (amiodarone)
  - o Vascor® (bepridil)
  - o Tambocor® (flecainide)
  - o Rythmol® (propafenone)
  - o Quinaglute dura® (quinidine)

## What should I tell my doctor before I take APTIVUS?

## Tell your doctor about all of your medical conditions, including if you:

- have liver problems or are infected with Hepatitis B or Hepatitis C. These patients may have worsening of their liver disease.
- are allergic to sulfa medicines.
- have hemophilia. APTIVUS may cause increased bleeding.
- have diabetes. APTIVUS may worsen your diabetes or high blood sugar levels.

- are pregnant or planning to become pregnant. It is not known if APTIVUS can harm your unborn baby. You and your doctor will need to decide if APTIVUS is right for you. If you take APTIVUS while you are pregnant, talk to your doctor about how you can be in the Antiretroviral Pregnancy Registry.
- are breast-feeding. Do not breast-feed if you are taking APTIVUS. You should not breast-feed if you have HIV because of the chance of passing the HIV virus to your baby. Talk with your doctor about the best way to feed your baby.
- are using estrogens for birth control or hormone replacement. Women who use estrogens for birth control or hormone replacement have an increased chance of developing a skin rash while taking APTIVUS. If a rash occurs, it is usually mild to moderate, but you should talk to your doctor as you may need to temporarily stop taking either APTIVUS or the other medicine that contains estrogen or female hormones.

Tell your doctor about all the medicines you take including prescription and nonprescription medicines, vitamins and herbal supplements. APTIVUS and many other medicines can interact. Sometimes serious side effects will happen if APTIVUS is taken with certain other medicines (see "Who should not take APTIVUS?").

- Some medicines cannot be taken at all with APTIVUS
- Some medicines will require a change in dosage if taken with APTIVUS
- Some medicines will require close monitoring if taken with APTIVUS.

Women taking birth control pills need to use another birth control method. APTIVUS makes birth control pills work less well.

Know all the medicines you take and keep a list of them with you. Show this list to all your doctors and pharmacists anytime you get a new medicine you take. They will tell you if you can take these other medicines with APTIVUS. Do not start any new medicines while you are taking APTIVUS without first talking with your doctor or pharmacist. You can ask your doctor or pharmacist for a list of medicines that can interact with APTIVUS.

## How should I take APTIVUS?

Take APTIVUS exactly as your doctor has prescribed. You should check with your doctor or pharmacist if you are not sure. You must take APTIVUS at the same time as NORVIR<sup>®</sup> (ritonavir). The usual dose is 500 mg (two 250 mg capsules) of APTIVUS, together with 200 mg (two 100 mg capsules or 2.5 mL of solution) of NORVIR, twice per day. APTIVUS with NORVIR must be used together with other anti-HIV medicines.

APTIVUS comes in a capsule form and you should swallow APTIVUS capsules whole. Do not chew the capsules.

- Always take APTIVUS with food.
- Do not change your dose or stop taking APTIVUS without first talking with your doctor.
- If you take too much APTIVUS, call your doctor or poison control center right away.
- If you forget to take APTIVUS, take the next dose of APTIVUS, together with NORVIR® (ritonavir), as soon as possible. Do not take a double dose to make up for a missed dose.

- It is very important to take all your anti-HIV medicines as prescribed and at the right times of day. This can help your medicines work better. It also lowers the chance that your medicines will stop working to fight HIV (drug resistance).
- When your APTIVUS supply starts to run low, get more from your doctor or pharmacy. This is very important because the amount of virus in your blood may increase if the medicine is stopped for even a short period of time. The HIV virus may develop resistance to APTIVUS and become harder to treat. You should NEVER stop taking APTIVUS or your other HIV medicines without talking with your doctor.

## What are the possible side effects of APTIVUS?

## APTIVUS may cause serious side effects, including:

- liver problems, including liver failure and death. Your doctor should do blood tests to monitor your liver function during treatment with APTIVUS. Patients with liver diseases such as Hepatitis B and Hepatitis C may have worsening of their liver disease with APTIVUS and should have more frequent monitoring blood tests.
- rash. Mild to moderate rash, including flat or raised rashes or sensitivity to the sun, have been reported in approximately 10% of subjects receiving APTIVUS. Some patients who developed rash also had joint pain or stiffness, throat tightness, or generalized itching.
- increased bleeding in patients with hemophilia. This can happen in patients taking APTIVUS or other protease inhibitor medicines.
- diabetes and high blood sugar (hyperglycemia). This can happen in patients taking APTIVUS or other protease inhibitor medicines. Some patients have diabetes before starting treatment with APTIVUS which gets worse. Some patients get diabetes during treatment with APTIVUS. Some patients will need changes in their diabetes medicine. Some patients will need new diabetes medicine.
- increased blood fat (lipid) levels. Your doctor should do blood tests to monitor your blood fat (triglycerides and cholesterol) during treatment with APTIVUS. Some patients taking APTIVUS have large increases in triglycerides and cholesterol. The long-term chance of having a heart attack or stroke due to increases in blood fats caused by APTIVUS is not known at this time.
- changes in body fat. These changes have happened in patients taking APTIVUS. and other anti-HIV medicines. The changes may include an increased amount of fat in the upper back and neck ("buffalo hump"), breast, and around the back, chest, and stomach area. Loss of fat from the legs, arms, and face may also happen. The cause and long-term health effects of these conditions are not known.

The most common side effects include diarrhea, nausea, vomiting, stomach pain, tiredness and headache. Women taking birth control pills may get a skin rash.

It may be hard to tell the difference between side effects caused by APTIVUS, by the other medicines you are also taking, or by the complications of HIV infection. For this reason it is very important that

you tell your doctor about any changes in your health. You should report any new or continuing symptoms to your doctor right away. Your doctor may be able to help you manage these side effects.

The list of side effects is not complete. Ask your doctor or pharmacist for more information.

#### **How should I store APTIVUS?**

- Store APTIVUS capsules in a refrigerator at approximately 36°F to 46°F (2°C to 8°C). Once the bottle is opened, the contents must be used within 60 days. Patients may take the bottle with them for use away from home so long as the bottle remains at a temperature of approximately 59°F to 86°F (15°C to 30°C). You can write the date of opening the bottle on the label. Do not use after the expiration date written on the bottle.
- Keep APTIVUS and all medicines out of the reach of children.

#### General advice about APTIVUS

Medicines are sometimes prescribed for purposes other than those listed in a Patient Information leaflet. Do not use APTIVUS for a condition for which it was not prescribed. Do not give APTIVUS to other people, even if they have the same condition you have. It may harm them.

This leaflet summarizes the most important information about APTIVUS. If you would like more information, talk with your doctor. You can ask your pharmacist or doctor for information about APTIVUS that is written for health professionals.

For additional information, you may also call Boehringer Ingelheim Pharmaceuticals, Inc. at 1-800-542-6257, or (TTY) 1-800-459-9906. You may also request information through the company website at <a href="http://us.boehringer-ingelheim.com">http://us.boehringer-ingelheim.com</a>.

## What are the ingredients in APTIVUS?

Active Ingredient: tipranavir

Major Inactive Ingredients: dehydrated alcohol, polyoxyl 35 castor oil, propylene glycol, mono/diglycerides of caprylic/capric acid and gelatin.

## Rx only

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APTIVUS Capsules are covered by U.S. Patents 5,852,195, 6,147,095, 6,169,181 and 6,231,887

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EXHIBIT B COPY OF U.S. PATENT 5,852,195



## United States Patent [19]

#### Romines et al.

## [11] Patent Number:

5,852,195

## [45] Date of Patent:

## Dec. 22, 1998

# [54] PYRANONE COMPOUNDS USEFUL TO TREAT RETROVIRAL INFECTIONS

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[21] Appl. No.: 809,224

[22] PCT Filed: May 4, 1995

[86] PCT No.: PCT/US95/05219

§ 371 Date: Nov. 4, 1996

§ 102(e) Date: Nov. 4, 1996

[87] PCT Pub. No.: WO95/30670

PCT Pub. Date: Nov. 16, 1995

## Related U.S. Application Data

[63]	Continuation of Ser. No. 349,361, Dec. 2, 1994, abandoned,
	which is a continuation of Ser. No. 238,817, May 6, 1994,
	abandoned.

[52] U.S. Cl. ...... 546/282.1; 546/162; 544/264; 544/286; 544/316; 548/311.1; 549/292

[58] Field of Search ...... 546/282.1

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Primary Examiner—Alan L. Rotman
Attorney, Agent, or Firm—Martha A. Gammill; Lawrence T.
Welch

## [57] ABSTRACT

The present invention relates to compounds of formulae (I) and (II) which are pyran-2-ones, 5,6-dihydro-pyran-2-ones, 4-hydroxy-benzopyran-2-ones, 4-hydroxy-cycloalkyl[b] pyran-2-ones, and derivatives thereof, useful for inhibiting a retrovirus in a mammalian cell infected with said retrovirus, wherein  $R_{10}$  and  $R_{20}$  taken together are formulae (III) and (IV).

$$\begin{array}{c}
R_{5} & OH \\
R_{1} & R_{3} \\
R_{2} & O
\end{array}$$
(I)

$$R_{10} \longrightarrow R_3$$

$$R_{10} \longrightarrow R_3$$

$$R_{10} \longrightarrow R_3$$

$$R_1$$
 (III)

$$(CH_2)_p$$
 (IV)

## 5 Claims, No Drawings

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# PYRANONE COMPOUNDS USEFUL TO TREAT RETROVIRAL INFECTIONS

#### **CROSS-REFERENCE**

This application is a 371 of PCT/US95/05219 filed May 4, 1995, which is a Continuation of U.S. Ser. No. 08/349,361 filed Dec. 2, 1994 (now abandoned), which is a Continuation of U.S. Ser. No. 08/238,817 filed May 6, 1994 (Abandoned).

The present invention relates to compounds useful for inhibiting a retrovirus in a human cell infected with said retrovirus. More particularly, the present invention provides pyran-2-ones, 5,6-dihydropyran-2-ones, 4-hydroxy-benzopyran-2-ones, 4-hydroxy-cycloalkyl[b]pyran-2-ones, and derivatives thereof as HIV-proteinase inhibitors.

#### BACKGROUND OF THE INVENTION

During the past decade, acquired immunodeficiency syndrome (AIDS) has progressed from having the status of a medical curiosity afflicting only a small number of individuals to a problem of major proportions, both medically and economically. John Saunders and Richard Storer, "New Developments in RT Inhibitors," DN&P 5(3), April 1992, pages 153–169. WHO figures reveal that more than 360,000 cases of AIDS have been reported worldwide, including nearly 175,000 cases in the U.S.A. Of these, approximately 100,000 worldwide (50,000 in the U.S.A.) were reported in the preceding 12-month period. In the U.S.A., the number of seropositive individuals is thought to be approximately two million, and estimates suggest that 5–10 million people worldwide may be seropositive. Saunders and Storer, page 153.

Since the first description of the malady in the early part of this decade, acquired immunodeficiency disease syndrome (AIDS) and its devastating consequences have been subjects of continuous and intense coverage in both the lay and scientific press. Indeed, an edition of Scientific American was entirely devoted to AIDS (Scientific American 289, #4 (1988)), and the literature on the disease and the virus is already so vast as to defy thorough citation.

On Mar. 20, 1987, the FDA approved the use of the compound, zidovudine (AZT), to treat AIDS patients with a recent initial episode of pneumocystis carinii pneumonia, AIDS patients with conditions other than pneumocystis carinii pneumonia or patients infected with the virus with an absolute CD4 lymphocyte count of less than 200/mm³ in the peripheral blood. AZT is a known inhibitor of viral reverse transcriptase, an enzyme necessary for human immunodeficiency virus replication. U.S. Pat. No. 4,724,232 claims a method of treating humans having acquired immunodeficiency syndrome utilizing 3'-azido-3'-deoxy-thymidine (azidothymidine, AZT).

Following the discovery of the anti-HIV activity of AZT, much effort has been focused on a wide variety of other dideoxynucleoside analogues in the search for superior agents. In the case of the 2'.3'-dideoxy series, ddC and ddI have shown potent activity against HIV in vitro and have been evaluated in clinical trials. Saunders and Storer, page 160. The compound ddC is currently being developed by Hoffman-La Roche Co. as a potential anti-AIDS drug. Its limiting toxicity in humans is peripheral neuropathy which is reversible at low doses. Raymond R. Schinazi, Jan R. Mead and Paul M. Feorino, "Insights Into HIV Chemotherapy," AIDS Research and Human Retroviruses, Vol. 8, Number 6, 1992, pages 963–990. It has been 65 approved by the FDA for AIDS therapy in combination with AZT. The compound ddI has also been evaluated in clinical

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trials. Its limiting toxicities are peripheral neuropathy and pancreatitis. It has also been shown to stimulate hepatic glycolysis leading to irreversible liver damage. Schinazi, Mead and Feorino, page 966. It has recently been approved by the FDA for the treatment of HIV-1 infections in adults and pediatric patients who are intolerant to or whose health has significantly deteriorated while on AZT treatment. Schinazi, Mead and Feorino, page 966.

Among these approved drugs, AZT is currently the only <sup>10</sup> drug that has been shown to decrease the mortality and frequency of opportunistic infections associated with AIDS. Schinazi, Mead and Feorino, page 963.

Human immunodeficiency virus (HIV) has long been recognized as the causative agent in AIDS, although a minority opinion to the contrary has been expressed (e.g., P. Duesberg, Proc. Natl. Acad. Sci., USA, 86:755-764 (1989)). Sequence analysis of the complete genomes from several infective and non-infective HIV-isolates has shed considerable light on the make-up of the virus and the types of molecules that are essential for its replication and maturation to an infective species. The HIV protease is essential for the processing of the viral gag and gag-pol polypeptides into mature virion proteins. L. Ratner, et al., Nature, 313:277-284 (1985); L. H. Pearl and W. R. Taylor, Nature, 329:351 (1987). HIV exhibits the same gag/pol/env organization seen in other retroviruses. L. Ratner, et al., above; S. Wain-Hobson, et al., Cell, 40:9-17 (1985); R. Sanchez-Pescador, et al., Science, 227:484-492 (1985); and M. A. Muesing, et al., Nature, 313:450-458 (1985).

Reverse transcriptase (RT) is an enzyme unique to retroviruses that catalyzes the conversion of viral RNA into double stranded DNA. Blockage at any point during the transcription process, by AZT or any other aberrant deoxynucleoside triphosphate incapable of elongation, should have dramatic consequences relative to viral replication. Much work on the RT target is in progress based, in large measure, upon the fact that nucleosides like AZT are easily delivered to cells. However, the inefficiency of phosphorylation steps to the triphosphate, and the lack of specificity and consequent toxicity, constitute major drawbacks to use of AZT and similar nucleosides having a blocked, or missing, 3'hydroxyl group.

The T4 cell receptor for HIV, the so-called CD4 molecule, has also been targeted as an intervention point in AIDS therapy. R. A. Fisher, et al., Nature, 331:76-78 (1988); R. E. Hussey, et al., Nature, 331:78-81 (1988); and K. C. Deen, et al., Nature, 331:82-84 (1988). The exterior portion of this transmembrane protein, a molecule of 371 amino acids (sCD4) has been expressed in Chinese hamster ovary (CHO) cells and Genentech (D. H. Smith, et al., Science, 238:1704-1707 (1987)) has had a product in clinical trials since the fall of 1987. CD4 has been shown to have a narrow spectrum of activity against wild-type virus and so far has failed to control HIV infection in humans. Schinazi, Mead and Feorino, page 963. The idea behind CD4 based therapy is that the molecules can neutralize HIV by interfering with viral attachment to T4, and other cells which express CD4 on their surfaces. A variant on this theme is to attach cell toxins to CD4 for specific binding and delivery to infected cells which display glycoprotein gp-120 on their surfaces. M. A. Till, et al., Science, 242:1166-1168 (1988); and V. K. Chaudhary, et al., Nature, 335:369-372 (1988).

Another therapeutic target in AIDS involves inhibition of the viral protease (or proteinase) that is essential for processing HIV-fusion polypeptide precursors. In HIV and several other retroviruses, the proteolytic maturation of the

gag and gag/pol fusion polypeptides (a process indispensable for generation of infective viral particles) has been shown to be mediated by a protease that is, itself, encoded by the pol region of the viral genome. Y. Yoshinaka, et al., Proc. Natl. Acad. Sci. USA, 82:1618–1622 (1985); Y. Yoshinaka, et al., J. Virol., 55:870–873 (1985); Y. Yoshinaka, et al., J. Virol., 57:826–832 (1986); and K. von der Helm, Proc. Natl. Acad. Sci., USA, 74:911–915 (1977). Inhibition of the protease has been shown to inhibit the processing of the HIV p55 in mammalian cell and HIV 10 replication in T lymphocytes. T. J. McQuade, et al., Science, 247:454 (1990).

The protease (or proteinase), consisting of only 99 amino acids, is among the smallest enzymes known, and its demonstrated homology to aspartyl proteases such as pepsin and 15 renin (L. H. Pearl and W. R. Taylor, Nature, 329:351-354 (1987); and I. Katoh, et al., Nature, 329:654-656 (1987)), led to inferences regarding the three-dimensional structure and mechanism of the enzyme (L. H. Pearl and W. R. Taylor, above) that have since been borne out experimentally. Active 20 HIV protease has been expressed in bacteria (see, e.g., P. L. Darke, et al., J. Biol. Chem., 264:2307-2312 (1989)) and chemically synthesized (J. Schneider and S. B. Kent, Cell, 54:363-368 (1988); and R. F. Nutt, et al., Proc. Natl. Acad. Sci., USA, 85:7129-7133 (1988)). Site directed mutagenesis 25 (P. L. Darke, et al., above); and N. E. Kohl, et al., Proc. Natl. Acad. Sci., USA, 85:4686-4690 (1988)) and pepstatin inhibition (P. L. Darke, et al., J. Biol. Chem., 264:2307-2312 (1989); S. Seelmeier, et al., Proc. Natl. Acad. Sci., USA, 85:6612-6616 (1988); C.-Z. Giam and I. Borsos, J. Biol. 30 Chem., 263:14617-14720 (1988); and J. Hansen, et al., EMBO J., 7:1785-1791 (1988)) have provided evidence for HIV protease's mechanistic function as an aspartyl protease. A study has demonstrated that the protease cleaves at the sites expected in peptides modeled after the regions actually 35 cleaved by the enzyme in the gag and pol precursor proteins during viral maturation. P. L. Darke, et al., Biochem. Biophys. Res. Communs., 156:297-303 (1988). X-ray crystallographic analysis of the HIV-protease (M. A. Navia, et al., Nature, 337:615-620 (1989)) and a related retroviral 40 enzyme from Rous sarcoma virus (M. Miller, et al., Nature, 337:576-579 (1989)) reveal an active site in the protease dimer that is identical to that seen in other aspartyl proteases, thus supporting the supposition (L. H. Pearl and W. R. Taylor, above) that the HIV enzyme is active as a dimer. See: 45 also Joseph A. Martin, "Recent Advances in the Design of HIV Proteinase Inhibitors," Antiviral Research, 17 (1992) 265-278.

To date, the scientific search for a fully effective and safe means of inhibiting retroviruses in a human hosting such a 50 virus, and thereby effectively treating diseases caused by such a virus, such as acquired immunodeficiency syndrome (AIDS), continues.

#### INFORMATION DISCLOSURE

JO 3227-923-A (Sawai Seiyaku KK) discloses the use of 4-hydroxy-coumarins as therapeutic agents for HIV-infected patients; however, unsubstituted 4-hydroxy-coumarin is the only compound specifically disclosed for this use.

WO 91/04663 (Univ. of Calif. at Oakland) discloses 60 6-amino-1,2-benzopyrones which are useful for treating viral diseases.

WO 91/12804 (Kabi Pharmaceutical), published 5 Sep. 1991, discloses the use of N-phenyl-N-methyl-1,2-dihydro-4-hydroxy-1-methyl-2-oxo-quinoline-3-carboxamide also 65 known as Linomide®, for the treatment of retrovirus infections.

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International Publication No. WO 89/07939, published 8 Sep. 1989, discloses specific coumarin compounds which are reverse transcriptase inhibitors.

U.S. Pat. Nos. 3,489,774 and 3,493,586 disclose 3-(beta-aryl-beta-(arylthio) (or aryl seleno) propionyl-coumarin and pyrone products useful as parasiticides.

Biochemical and Biophysical Research Communications, Vol. 188, No. 2, 1992, pages 631–637, discloses chromones bearing hydroxyl substituents and a phenolic group at the 2-position (flavones) as having anti-HIV-1 proteinase activity

Antimicrobial Patent Fast-Alert, Week Ending 4 Sep. 1992, disclose gamma-pyrones, gamma-pyridones and gamma-thio-pyrones as antiviral agents.

International Publication Nos. WO 92/04326, 92/04327 and 92/04328, all published 19 Mar. 1992, disclose antiviral heterocyclic derivatives, such as quinolinones and benzopyranones, as replication inhibitors for treating herpes simples 1 and 2, cytomegalovius and Epstein-Barr virus.

C.A. Selects: Antitumor Agents, Issue 19, 1992, page 25, No. 117:90147q (PCT International Application WO 92 06,687) discloses the preparation of 5-iodo-5-amino-1,2-benzopyrones and analogs as cytostatic and antiviral agents.

Nowhere do these references teach or suggest the use of 4-hydroxy- $\alpha$ -pyrones as HIV protease inhibitors or as having antiviral activity.

Phytochemistry, 31(3):953-956 (1992), discloses compounds, such as 4-hydroxy-α-(4-methoxyphenyl)-6-[2-(4-methoxyphenyl)ethenyl]-2-oxo-, methyl ester, (E)-(-)-2H-pyran-3-acetic acid.

Tetrahedron, 48(9):1695–1706 (1992), (see also Tetrahedron Lett., 30(23):3109–12 (1989)), discloses compounds, such as 3-[1-(4-chlorophenyl)-3-(4-nitrophenyl)-2-propenyl]-4-hydroxy-6-methyl-2H-pyran-2-one; 3-[3-(4-chlorophenyl)-1-(4-nitrophenyl)-2-propenyl]-4-hydroxy-6-methyl-2H-pyran-2-one; 4-hydroxy-3-[3-(4-methoxyphenyl)-1-(4-nitrophenyl)-2-propenyl]-6-methyl-2H-pyran-2-one; and 4-hydroxy-3-[1-(4-methoxyphenyl)-3-(4-nitrophenyl)-2-propenyl]-6-methyl-2H-pyran-2-one.

Tennen Yuki Kagobutsu Toronkai Koen Yoshishu, 30:17-24 (1988), discloses compounds, such as 4-hydroxy-β-(4-methoxyphenyl)-6-[2-(4-methoxyphenyl)ethenyl]-2-oxo-, methyl ester, (E)-(-)-2H-pyran-3-propanoic acid.

Chem. Absts. 53:15072f discloses compounds, such as  $\alpha$ -1,3-dihydroxy-2-butenylidene- $\beta$ -ethyl-,  $\delta$ -lactone, hydrocinnamic acid.

Chem. Absts. 53:15072c discloses compounds, such as  $\alpha$ -1,3-dihydroxy-2-butenylidene- $\beta$ -isopropyl-,  $\delta$ -lactone, hydrocinnamic acid.

Arch. Pharm. (Weinheim, Ger.), 316(12):988-94 (1983), discloses compounds, such as 3-[1-(4-chlorophenyl)-3-oxobutyl]-4-hydroxy-6-methyl-2H-pyran-2-one; and 3-[1-(4-chlorophenyl)propyl]-4-hydroxy-6-methyl-2H-pyran-2-one.

Chem. Ber., 110(3):1047-57 (1977), discloses compounds, such as 6-(3,4-dimethoxyphenyl)-3-[2-(3,4-dimethoxy-phenyl)-1-(4-methoxy-2-oxo-2H-pyran-6-yl) ethyl]-4-hydroxy-2H-pyran-2-one; and 3-[2-(3,4-dimethoxyphenyl)-1-(4-methoxy-2-oxo-2H-pyran-6-yl) ethyl-4-hydroxy-6-[2-(4-methoxyphenyl)ethyl]-2H-pyran-2-one.

J. Heterocycl. Chem., 23(2):413-16 (1986), discloses compounds, such as 3-[(4-chlorophenyl)-1-piperidinylmethyl]-4-hydroxy-6-methyl-2H-pyran-2-one.

The following published PCT applications disclose peptides useful as retroviral protease inhibitors: International

Publication No. WO 91/06561, published 16 May 1991; and International Publication No. WO 92/17490, published 15 Oct. 1992.

The following references disclose pyrone compounds which are believed to be representative of those known in 5 the art:

EP-443449 (German language) discloses 3-hexyl-5,6dihydro-6-pentyl-2H-pyran-2-one and 3-ethyl-6-hexadecyl-5,6-dihydro-4-hydroxy-2H-pyran-2-one. Pestic. Sci., 27(1): 45-63 (1989), discloses 5,6-dihydro-4-hydroxy-6-methyl-6- 10 (1-methyl-1-propenyl)-3-(1-oxobutyl)-2H-pyran-2-one; and 6-cyclopropyl-5,6-dihydro-4-hydroxy-6-methyl-3-(1oxobutyl)-2H-pyran-2-one. Acta. Chem. Scand., 43(2): 193-95 (1989), discloses 4-(acetyloxy)-5,6-dihydro-3,6dimethyl-2H-pyran-2-one. J. Org. Chem., 54(14):3383-9 15 (1989), discloses 5,6-dihydro-4-hydroxy-3,6,6-trimethyl-2H-pyran-2-one. J. Org. Chem., 53(6):1218-21 (1988); and Tetrahedron Lett., 34(2):277-80 (1993), discloses 3-hexyldihydro-6-undecyl-2H-pyran-2,4 (3H)-dione, (6R)-. J. Chem. Soc. Perkins Trans., 1(6):1157-9 (1985), discloses <sup>20</sup> dihydro-3-methyl-6-nonyl-6-[[(tetrahydro-2H-pyran-2-yl) oxy]methyl]-2H-pyran-2,4 (3H)-dione. J. Chem. Ecol., 9(6) :703-14 (1983), discloses 5,6-dihydro-4-hydroxy-3,6dimethyl-2H-pyran-2-one. J. Org. Chem., 48(7):1123-5 (1983), discloses 6-(2-chloro-1-methylethenyl-5,6-dihydro- 25 4-hydroxy-3-methyl-2H-pyran-2-one, (Z)-(.+-.)-. Acta. Chem. Scand., 43(2):193-95 (1989); and Tetrahedron Lett., 21(6):551-4 (1980), discloses 5,6-dihydro-4-hydroxy-3,6dimethyl-2H-pyran-2-one. Helv. Chem. Acta, 59(7) methyl-6-oxo-2H-pyran-2-yl)methyl]-2,6-piperidinedione. Acta. Chem. Scand., 30(7):613-18 (1976); and Tetrahedron Lett., 22:1903-4 (1976), discloses 5,6-dihydro-4-hydroxy-3-methyl-6-(1-methyl-1-propenyl)-2H-pyran-2-one, (E)-. 3,3'-[(4-nitrophenyl)methylene]bis[5,6-dihydro-4-hydroxy-6-methyl-2H-pyran-2-one; and 3,3'-(phenylmethylene)bis [5,6-dihydro-4-hydroxy-6-methyl-2H-pyran-2-one are disclosed in Synth. Commun., 20(18):2827-2836, 1990.

WO 93/07868, published 29 Apr. 1993, discloses new nitroso-benzopyrone, -benzamide and -isoquinolinone derivatives as adenosine di-phospho:ribose transferase inhibitors for treating viral infections and cancer.

WO 93/07128, published 15 Apr. 1993, relates to substituted cyclic carbonyls and derivatives thereof useful as 45 retroviral protease inhibitors.

J. Indian Chem. Soc., 69:397-398 (July 1992), discloses that coumarin-4-acetic acids were screen for their anticancer and anti-AIDS activities and were found to be inactive.

The Journal of Antibiotics, 46(7):1126 (July 1993), dis-50 closes germicidin, which is 6-(2-butyl)-3-ethyl-4-hydroxy-2-pyrone, to be an autoregulative germination inhibitor of Streptomyces viridochromogenes NRRL B-1551.

Derwent Abstracts, 93-168920/21 of EP 543201 discloses the use of coumarin derivatives, such as 1-(N-morpholyl)-6-(4-hydroxybenzoic acid ethyl ester) hexane, for the treatment of viral infections, such as influenza or acute rhinitis.

- J. Org. Chem., 48(22):3945-7 (1983); and Chem. Pharm. Bull., 29(10):2762-8 (1981); disclose compounds such as 60 4-hydroxy-6-(3-pyridinyl)-2H-pyran-2-one.
- J. Labelled Compd. Radiopharm., 28(10):1143-8 (1990), discloses compounds such as 4-hydroxy-6-methyl-2Hpyran-2-one.
- J. Am. Chem. Soc., 113(25):9585-95 (1991), discloses 65 compounds such as 3-(3-phenyl-2-propen-1-yl)-6-methyl-4hydroxy-2H-pyran-2-one.

CA 54:14239d and CA 53:4272c disclose compounds such as  $\alpha$ -( $\alpha$ , $\gamma$ -dihydroxycinnamylidene)-,  $\delta$ -lactone hydrocinnamic acid.

CA 53:15072f discloses compounds such as  $\alpha$ -1,3dihydroxy-2-butenylidene-β-ethyl-, δ-lactone hydrocinnamic acid.

Synth. Commun., 20(18):2827-36 (1990), discloses compounds such as 3,3'-[(4-nitrophenyl)methylene]bis[5,6dihydro-4-hydroxy-6-methyl-2H-pyran-2-one, and 3,3'-(phenylmethylene)bis[5,6-dihydro-4-hydroxy-6-methyl-2H-pyran-2-one.

J. Org. Chem., 54(14):3383-9 (1989), discloses compounds such as 5,6-dihydro-4-hydroxy-3,6,6-trimethyl-2Hpyran-2-one.

Derwent Abstract, 92-166863/20, of EP 553248 discloses new optionally substituted 5-iodo-6-amino-1,2benzopyrone derivatives, which are adenosine di:phosphoribose inhibitors, for treatment and prevention of viruses and tumors associated with AIDS.

Synthesis of Heterocycles. XV. 4-Hydroxy-2pyronocyclenes. E. Ziegler, H. Junek, and E. Nolken, Monatsh., 89:678-82 (1958) (CA 53:12283-4) discloses compounds such as the following: 4-hydroxy-3-benzyl-5,6octamethylene-2-pyrone; 4-hydroxy-3-benzyl-5,6pentamethylene-2-pyrone; 4-hydroxy-3-benzyl-5,6heptamethylene-2-pyrone; 4-hydroxy-3-benzyl-5,6hexamethylene-2-pyrone; and 4-hydroxy-3-benzyl-5,6tridecamethylene-2-pyrone.

R. Effenberberger, T. Ziegler, K.-H. Schonwalder, T. :2393-2401 (1976), discloses 4-[(3,6-dihydro-4-hydroxy-5-30 Kesmarszky, B. Bauer, Chem. Ber 119:3394-3404 (1986), discloses pyrone intermediates, such as those of formula J-1 (wherein n is 4; refer to Chart J below).

> Monatsh. Chem., 119(6-7):727-37 (1988) (CA 110(13): 114430k) discloses the compounds 8H-acenaphtho[1,2-b] pyran-8-one, 10-hydroxy-9-(phenylmethyl)-; and indeno[2, 1-b]pyran-3(5H)-one, 1-hydroxy-2-(phenylmethyl)-.

CA 54:14239b discloses the compound 3-benzyl-4hydroxy-2-oxoindeno-[1,2-b]pyran.

Monatsh. Chem., 113(4):475-84 (1982) discloses compounds such as 6,7-dihydro-4-hydroxy-6-(3-methylphenyl)-7-phenyl-3-(phenylmethyl)-pyrano[2,3-c]pyrrole-2,5-dione; and 6,7-dihydro-4-hydroxy-6,7-diphenyl-3-(phenylmethyl)pyrano[2,3-c]pyrrole-2,5-dione.

Monatsh. Chem. 90:594-9 (1959) (CA 54:14238g,h) discloses compounds such as 5H-benzocycloheptene-8-acrylic acid, α-benzyl-6,7-dihydro-β,-9-dihydroxy-,δ-lactone; and 3-benzyl-5,6,7,8-tetrahydro-4-hydroxy-8-isopropyl-5methyl-coumarin.

Bull. Soc. Chim. Fr. 5:1719-23 (1069) (Fr) (CA 71(21): 101655p) discloses the compound 3-benzyl-5,6,7,8tetrahydro-4-hydroxy-coumarin.

WO 8804652 (equivalent AU 8810440 (Jap.)) discloses the compound 3-(4-chloro-2-nitrobenzoyl)-5,6,7,8tetrahydro-4-hydroxy-2H-1-benzopyran-2-one.

Monatsh. 92:246-53 (1961) (Gr) (CA 55:27296d) discloses the compound 3-(3,5-dimethylsalicyl)-5,6,7,8tetrahydro-4-hydroxy-coumarin.

CA 94(9):65472r discloses 5,6,7,8-hexahydro-3-phenyl-2-H-cycloocta[b]pyran-2-one; and 6,7,8,9-tetrahydro-4hydroxy-3-phenyl-cyclohepta[b]pyran-2(5H)-one.

J. Org. Chem. 28(11):3112-14 (1963) (CA 59:15185e) discloses the compound hexanedioic acid, 2-[hydroxy(2hydroxy-1-cyclopenten-1-yl)methylene]-,  $\delta$ -lactone.

Antimicrobial Patent Fast-Alert, Week Ending 30 Apr. 1993, discloses cyclic ureas and analogues useful as retroviral protease inhibitors.

Many 4-hydroxy-coumarin type compounds are known. For example, these references—CA 54:577e,g,h (1960); U.S. Pat. No. 2,872,457 (CA 53:12305e (1959)); CA 51:14826f,h (1957); U.S. Pat. No. 2,723,276 (CA 52:5480g,h (1958)); CA 51:14827a,b (1957); CA 51:16453a (1957); CA 54:5699d (1960); CA 54:16450f (1960); CA 53:22454a (1959); and CA 53:20046a—disclose compounds such as the following: 4-hydroxy-3-(1-phenylbutyl)coumarin; 4-hydroxy-3-(1-phenylpentyl)-coumarin; 3-(cyclohexylphenylmethyl)-4-hydroxycoumarin; 4-hydroxy-3-(2-methyl-1-phenylpropyl)-coumarin; 4-hydroxy-3-(2-phenylpropyl) coumarin; 4-hydroxy-3-(1,3diphenylpropyl)-coumarin; 4-hydroxy-3-(1-(4methylphenyl)-butyl)-coumarin; 4-hydroxy-3-(1-(1naphthyl)-propyl)-coumarin; 4-hydroxy-7-methyl-3-(1phenylpropyl)-coumarin; 7-chloro-4-hydroxy-3-(1-15 phenylpropyl)-coumarin; 4-hydroxy-3-[1-(4methoxyphenyl)propyl]-coumarin; 3-(.alpha.-ethyl-pfluorobenzyl)-4-hydroxy-coumarin; 3-(a-ethyl-pmethoxybenzyl)-4-hydroxy-coumarin; and 3-(1-phenylpropenyl)-4-hydroxy-coumarin.

To the best of our knowledge, from our review, these references do not disclose the use of these compounds as HIV protease inhibitors. They are disclosed as being useful as: rodenticides, lowering the prothrombin level of the blood, blood anticoagulants, and pesticides.

Additional 4-hydroxy-coumarin compounds with similar uses have been disclosed in the following references:

Indian J. Chem., Sect. B, 25B: 1167–70 (1986) (CA 107(17):154201f) and CA 93(23):220546t discloses the compound 4-Hydroxy-3-(1-phenyl-2-propenyl)-coumarin.

CA 96(19):157432x; CA 90(1):1707f; CA 84(9):55338f; CA 79(13):74969a; and CA 71(15):69677j disclose the compound 4-hydroxy-3-[1-(1,2,3,4-tetrahydro)naphthyl]-coumarin; CA 54:579e discloses the compound 4-hydroxy-3-[1-indanyl]-coumarin; CA 63:14743c discloses the compound 4-hydroxy-3-(1-naphthylmethyl)-coumarin; CA 63:5589c discloses the compound 3-(1'-(2-methoxy,3-methyl,5-chloro-phenyl)propyl)-4-hydroxy-coumarin; CA 64:12969b discloses the compound 3-(α-acetonyl-α-acetylbenzyl)-4-hydroxy-coumarin.

CA 79(13):74969a; Chim. Ther. 7(4):300-6 (1972) (Fr) (CA 78(7):38016h); CA 52:5399b; CA 54:5699e; CA 54:579e; and CA 72(15):78882v disclose 4-hydroxycoumarin compounds substituted at the 6- or 7-position by, e.g., methyl, methoxy and chloro.

J. M. Mulder, U.S. Pat. No. 3,835,161, 10 Sep. 1974, discloses the compound 3-[1-[4-(2-bromoethyl)phenyl] ethyl]-4-hydroxy-2H-1-benzopyran-2-one.

Merck Index, Eleventh Edition, (1989), Entry 9950, discusses Warfarin, its chemical name—3-α-phenyl-β-50 acetylethyl-4-hydroxycoumarin—and its uses as a rodenticide and an anticoagulant. J. Med. Chem., 1978, Vol. 21, No. 2:231–234, discloses the antivitamin K activity of warfarin and discusses the anticoagulant activity of several 3-substituted 4-hydroxycoumarins such as 4-Hydroxy-3-(1-55 phenylbutyl)-coumarin; and 4-hydroxy-3-(α-methylbenzyl)-coumarin. J. Am. Chem. Soc. 83:2676–9 (1961) (CA 55:22306e (1961)) discusses the resolution and absolute configuration of warfarin and discloses the preparation of compounds such as 4-hydroxy-3-(1-phenylbutyl)-60 coumarin.

Journal of Labelled Compounds and Radiopharmaceuticals Vol. XXIII, No. 2:137-148 (1986), discloses several deuterium labelled metabolites of warfarin and phenprocoumon, such as the deuterium labelled analog of 65 the compound 4-hydroxy- 7-methoxy-3-(1-phenylpropyl)-coumarin.

J48023942 discloses compounds, such as 4-hydroxy-3-(α-methylbenzyl)-coumarin; 4-hydroxy-3-(3-methyl-1-phenylbutyl)-coumarin; and 2H-1-benzopyran-2-one, 4-hydroxy-7-methoxy-3-(1-phenylpropyl)-(also cited in preceding reference) and their use as rodenticides.

Tr. Voronezh. Teckhnol. Inst. 19(2):27-30 (1971), Abstract No. 1zh274 (Russian language), discloses the compound 4-hydroxy-3-phenethylcoumarin. This reference and Helv. Chim. Acta 74(7):1451-8 (1991) disclose the compound of 4-hydroxy-3-(3-phenylpropyl)coumarin.

J. Org. Chem. 33(1):437–8 (1968); and Eur. J. Med. Chem.—Chim Ther. 12(2):125–30 (1977) disclose compounds such as 4-hydroxy-3-diphenylmethylcoumarin.

U.S. Pat. No. 3,764,693 discloses the compound 4-hydroxy-3-(3-hydroxy-1-phenylbutyl)-coumarin and its anticoagulating and rodenticidal activity.

J. Med. Chem. 18(5):513–19 (1975) (CA 83(5):37913q);
J. Chromatogr. 338(2):325–34 (1985); J. Chromatogr. 562 (1-2):31–8 (1991); J. Labelled Compds. Radiopharm. 23(2): 137–48 (1986) (cited previously); and J. Chromatogr. 529 (2):479–85 (1990) disclose compounds such as 4-hydroxy-3-[1-[3-(phenylmethoxy)phenyl]propyl]-2H-1-benzopyran-2-one; 4-hydroxy-8-(phenylmethoxy)-3-(1-phenylpropyl)-2H-1-benzopyran-2-one; 4-hydroxy-6-methoxy-3-(1-phenylpropyl)-coumarin; 4-hydroxy-6-methoxy-3-(1-phenylpropyl)-coumarin; 4,6-dihydroxy-3-(1-phenylpropyl)-coumarin; 4,6-dihydroxy-3-[1-(3-hydroxyphenyl)propyl]-coumarin; 4-hydroxy-3-[1-(3-hydroxyphenyl)propyl]-coumarin; and p-chlorophenprocoumon.

AIDS 1993, Vol. 7, No. 1, pages 129-130, discusses the effect of warfarin on HIV-1 replication and spread.

CA Selects: AIDS & Related Immunodeficiencies, Issue 35 24, 1993, Abstract 119:195147j discloses the inhibitory effect of a single dose of coumarin derivatives, warfarin, 4-hydroxy-coumarin, umbelliferone, on HIV-1 replication and cell-mediated or cell-free viral transmission.

At the First National Conference on Human Retroviruses and Related Infections, 12–16 Dec. 1993, Washington, D.C., it was disclosed that coumarins, such as warfarin, and pyrones, such as 3-(thiophenyl)-6-phenyl-4-hydroxypyrone, displayed HIV protease inhibition in an assay.

Biochemical and Biophysical Research Communications, Vol. 201, No. 1, pages 290–294 (30 May 1994) discloses that warfarin and structurally related coumarin analogs are HIV-1 protease inhibitors.

J. Med. Chem. 37:2664-2677 (1994) discloses 4-hydroxy-3-(3-phenoxypropyl)-2H-1-benzopyran-2-one and structural analogs, especially 4,7-dihydroxy-3-[4-(2-methoxyphenyl)butyl]-2H-1-benzopyran-2-one, as HIV-1 protease inhibitors.

Biochemical and Biophysical Research Communications, Vol. 200, No. 3, pages 1658–1664 (16 May 1994) discloses that 4-hydroxy-3-(3-phenoxypropyl)-1-benzopyran-2-one and 4-hydroxy-6-phenyl-3-(phenylthio)-pyran-2-one, and structural analogs of these compounds, are inhibitors of HIV-1 protease.

J. Am. Chem. Soc. 116:6989-6990 (1994) discloses 4-hydroxy-6-phenyl-3-(phenylthio)pyran-2-one, and structural analogs thereof, are HIV-1 protease inhibitors.

Acta. Virol. 37:241-250 (1993) discloses the anti-HIV activity of coumarin derivatives, warfarin, 4-hydroxy-coumarin and umbelliferone.

Antiviral Research 24:275-288 (1994) discloses bicyclic imidazo derivatives (imidazothiazoles and

imidazopyridines) which inhibit HIV-1 through interaction with reverse transcriptase (RT).

U.S. Pat. No. 3,325,515 (J. Schmitt, et al.) discloses coumarin derivatives, such as 3-(4-hydroxy-3-coumarinyl)-3-phenyl-1-propionic acid methyl ester, as exhibiting anticoagulant activity.

U.S. Pat. No. 2,723,277 (A. Grussner, et al.) discloses malonic acid derivatives, such as 3-[1'-(p-chloro-phenyl)-propyl]-4-hydroxy-coumarin, as anti-coagulant agents.

FR, A, 1276654 discloses 4-hydroxy-coumarins, such as 10 (2'-hydroxy)-3-benzyl-4-hydroxycoumarin, which have anti-coagulant, anti-bacterial or anti-fungal properties.

BE, A, 674997 discloses 4-hydroxycoumarin derivatives, such as 3-(5-methoxytetralyl-(1))-4-hydroxycoumarin as agents for fighting rodents.

GB, A, 734142 discloses the preparation of 3-substituted-4-hydroxycoumarins, such as 3-(1-phenyl-2-acetyl)-ethyl-4-hydroxycoumarin and 3-(1-furan-2-acetyl)-ethyl-4-hydroxycoumarin, which are effective as anti-coagulants and rodenticides.

"The Application of Computer-Assisted Drug Design in the discovery of Nonpeptide HIV-1 Protease Inhibitors", Parke-Davis Pharm. Res., Keystone Symposia, 5–11 Mar. 1994, Santa Fe, N. Mex., discloses 4-hydroxy-3-(3-phenoxypropyl)-1-benzopyran-2-one as an HIV protease 25 inhibitor.

Structural Biology, 1(1):199-200 (April 1994) discloses that the rat poison warfarin was a useful lead in the search for HIV proteinase inhibitors.

CA 85:78002b (1976) discloses 3-(2,4,6-30 trihydroxybenzyl)-4-hydroxy-2H-pyran-2-one derivatives as having anti-bacterial activity.

FR, A, 1092278 (Hoffman) (1955) discloses the preparation of coumarin derivatives, such as 3-[1'-phenyl-propene-(1')-yl)-4-hydroxycoumarin.

International Publication No. WO 94/11361, published 26 May 1994, discloses pyran-2-ones and 5,6-dihydroxypyran-2-ones as retroviral protease inhibitors.

International Publication No. WO 94/18188, published 18 Aug. 1994, discloses 4-hydroxy-benzopyran-2-ones and 4-hydroxy-cycloalkyl[b]pyran-2-ones as retroviral protease inhibitors.

The following references were cited against the immediate parent application as disclosing the state of the art:

U.S. Pat. No. 3,651,091 (Boschetti, et al.); U.S. Pat. No. 4,262,013 (Mistui, et al.); U.S. Pat. No. 4,900,754 (Regan, et al.); U.S. Pat. No. 5,294,724 (Jendralla, et al.); Australian Patent Specification 219,371 (Enders, et al.); Canadian Patent No. 1,171,424 (Willard, et al.); British Patent Specification 836,740 (Bayer); European Patent Application 0 024 348 (Willard, et al.); European Patent Application 0 588 137 (Fischer, et al.); French Patent No. 1,276,654 (Molho) (cited above); and International Publication No. WO 94/1136 (Thaisrivongs, et al.) (cited above).

"Collaborative Structure-Based Design of Small Organic Molecules as Inhibitors of HIV Proteases," Keystone Symposia, Santa Fe, N. Mex. (5-11 Mar. 1994), discloses the crystallographic complexes of HIV-1 and HIV-2 protease with compounds, such as  $3-(\alpha-\text{ethylbenzyl})-6-(\alpha-60 \text{ ethylphenethyl})-4-hydroxy-2H-pyran-2-one.$ 

"Discovery and Properties of Small Organic Molecules Inhibiting HIV-1 Protease," Keystone Symposia, Santa Fe, N. Mex. (5–11 Mar. 1994), discloses an assay for determining inhibitory activity of compounds, such as 3-( $\alpha$ -65 ethylbenzyl)-6-( $\alpha$ -ethylphenethyl)-4-hydroxy-2H-pyran-2-one.

"Structure-based Design of Non-peptide HIV Protease Inhibitors," 35th Annual Buffalo Medicinal Chemistry Symposium, Buffalo, N.Y. (22–25 May 1994), discloses compounds, such as 3-( $\alpha$ -ethylbenzyl)-6-( $\alpha$ -ethylphenethyl)-4-hydroxy-2H-pyran-2-one, as potential anti-HIV therapeutic agents.

In Hruby et. al. (J. Org. Chem., 58 (26):7567 (1993), a description of the copper catalyzed addition of an aryl Grignard to an unsaturated chiral amide, 3-(2-butenoyl)-4phenyl-2-oxazolidinone, is given. In Evans et. al. (J. Am. Chem. Soc., 112:8215 (1990), the reaction between a chiral amide and 2-methoxy-2-methyl-1,3-dioxoline is described. The preparation of 2-methoxy-2-methyl-1,3-dioxoline is found in Santry et. al. (J. Am. Chem. Soc., 110 (9):2910 (1988). For references on the reaction between an ester enolate and a ketone, refer to Dongala et. al., Tetrahedron Letters, 4983 (1973), and Mitsui et. al., Tetrahedron, 23:4271 (1967). For references on the reaction between an amide enolate and a ketone, refer to Viteva et. al., Tetrahedron 50:7193 (1994); Oare et. al., J. Org. Chem. 55:132 (1990); Hullot et. al., Can. J. Chem. 55:266 (1977); Woodbury et. al., J. Org. Chem. 42:1688 (1977); Stefanovsky et. al., Tetrahedron 42:5355 (1986); and Mathew et. al., U.S. Pat. No. 5,284,975.

G. Carganico, P. Cozzi, G. Orsini, J. Med. Chem., 26:1767–1769 (1983), discloses synthesized compounds with a methyl and a hydroxyl group at the 4-position of the dihydropyrone ring and no substitution at the 3-position. The compounds of the present invention have a ketone at the 4-position (which may be in enol form) and substitution at the 3-position.

D. T. Witiak et al., J. Med. Chem., 31:1437–1445 (1988), discloses benzopyran-2-ones with a hydroxy group at the 3-position. The compounds of the present invention have alkyl substitution at that position.

B. Tait, Winter Conference on Bioorganic Medicinal Chemistry, 29 Jan.-2 Feb. 1995, Steamboat Springs, Colorado, disclosed a dihydropyrone having a phenyl group and a pentyl group at the 6-position and a —S—CH<sub>2</sub>—CH<sub>2</sub>-phenyl group at the 3-position in the HIV protease area.

J. V. N. Vara Prasad, et al., J. Med. Chem., 38:898-905 (1995), discloses 4-hydroxy-6-phenyl-2-oxo-2H-pyran-3-yl)thiomethanes, such as (+)-3-[cyclopentyl (cyclopentylthio)methyl]-4-hydroxy-6-phenyl-2H-pyran-2-one, as HIV-1 protease inhibitors.

#### SUMMARY OF THE INVENTION

The present invention provides:

A compound of the formula I

wherein R<sub>1</sub> is H-;

wherein R<sub>2</sub> is

a)  $C_3$ – $C_5$  alkyl,

b) phenyl-(CH<sub>2</sub>)<sub>2</sub>—,

c) het—SO<sub>2</sub>NH—(CH<sub>2</sub>)<sub>2</sub>—,

d) cyclopropyl-(CH<sub>2</sub>)<sub>2</sub>—,

e) F-phenyl-(CH<sub>2</sub>)<sub>2</sub>---,

f) het—SO<sub>2</sub>NH-phenyl-, or

g)  $F_3C-(CH_2)_2-$ ;

or wherein  $R_1$  and  $R_2$  taken together are a double bond; wherein  $R_3$  is the moiety of formula X wherein  $R_4$  is

- a) phenyl,
- b) het,
- c) cyclopropyl,
- d)  $H_3C-[O(CH_2)_2]_2-$
- e) het—SO<sub>2</sub>NH—,
- f) Br—,

```
g) N_3—, or
                                                                                               c) het—SO_2NH—(CH_2)_2—
    h) HO<sub>3</sub>S(CH<sub>2</sub>)<sub>2</sub>—N(CH<sub>3</sub>)—C(O)—(CH<sub>2</sub>)<sub>6</sub>—C(O)—
                                                                                           or wherein R<sub>1</sub> and R<sub>2</sub> taken together are a double bond;
                                                                                           wherein R<sub>3</sub> is the moiety of formula X
wherein R_5 is —H;
                                                                                           wherein R<sub>4</sub> is
                                                                                   5
wherein R<sub>6</sub> is
                                                                                              a) phenyl,
   a) R<sub>4</sub>—(CH<sub>2</sub>)<sub>n</sub>—CH(R<sub>8</sub>)—,
b) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—CH<sub>2</sub>—,
                                                                                               b) het,
                                                                                               c) cyclopropyl,
                                                                                              d) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—,
e) het—SO<sub>2</sub>NH—,
    c) C<sub>3</sub>-C<sub>5</sub> alkyl,
    d) phenyl-(CH<sub>2</sub>)<sub>2</sub>-
                                                                                  10
   e) het—SO<sub>2</sub>NH—(CH<sub>2</sub>)<sub>2</sub>—,
f) (HOCH<sub>2</sub>)<sub>3</sub>C—NH—C(O)—NH—(CH<sub>2</sub>)<sub>3</sub>—,
                                                                                              f) Br---,
                                                                                               g) N<sub>3</sub>---, or
                                                                                               h) HO_3S(CH_2)_2—N(CH_3)—C(O)—(CH_2)_6—C(O)—
    g) (HO_2C)(H_2N)CH - (CH_2)_2 - C(O) - NH
       (CH<sub>2</sub>)<sub>3</sub>-
                                                                                                  NH-;
    h) piperazin-1-yl-C(O)—NH—(CH<sub>2</sub>)<sub>3</sub>,
i) HO<sub>3</sub>S(CH<sub>2</sub>)<sub>2</sub>—N(CH<sub>3</sub>)—C(O)—(CH<sub>2</sub>)<sub>6</sub>—C(O)—
                                                                                           wherein R<sub>5</sub> is -H;
                                                                                           wherein R6 is
       NH-(CH<sub>2</sub>)<sub>3</sub>-
                                                                                              a) R<sub>4</sub>—(CH<sub>2</sub>)<sub>n</sub>—CH(R<sub>8</sub>)—,
b) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—CH<sub>2</sub>—
c) C<sub>3</sub>—C<sub>5</sub> alkyl,
   j) cyclopropyl-(CH<sub>2</sub>)<sub>2</sub>--,
   k) F-phenyl-(CH<sub>2</sub>)<sub>2</sub>-
   l) het-SO<sub>2</sub>NH-phenyl, or
                                                                                              d) phenyl-(CH<sub>2</sub>)<sub>2</sub>—,
e) het—SO<sub>2</sub>NH—(CH<sub>2</sub>)<sub>2</sub>—,
                                                                                  20
   m) F_3C—(CH_2)_2—;
wherein n is zero (0), one (1) or two (2);
                                                                                              f) (HOCH_2)_3C—NH—C(O)—NH—(CH_2)_3—,
                                                                                               g) (HO_2C)(H_2N)CH-(CH_2)_2-C(O)-NH-
wherein R<sub>7</sub> is
    a) cyclopropyl,
                                                                                               h) piperazin-1-yl-C(O)-NH-(CH<sub>2</sub>)<sub>3</sub>, or
   b) CH_3—CH_2—, or
                                                                                  25
   c) t-butyl;
                                                                                               i) HO_3S(CH_2)_2—N(CH_3)—C(O)—(CH_2)_6—C(O)—
                                                                                                  NH-(CH_2)_3-;
wherein R<sub>8</sub> is
   a) ---CH<sub>2</sub>---CH<sub>3</sub>, or
                                                                                           wherein n is zero (0), one (1) or two (2);
   b) —CH2-cyclopropyl;
                                                                                           wherein R<sub>7</sub> is
                                                                                               a) cyclopropyl,
wherein Ro is
                                                                                              b) CH<sub>3</sub>—CH<sub>2</sub>—, or
   a) -NR<sub>12</sub>SO<sub>2</sub>-het,
                                                                                              c) t-butyl;
   b) -NR<sub>12</sub>SO<sub>2</sub>-phenyl substituted by zero (0) or one (1)
                                                                                           wherein R<sub>8</sub> is
                                                                                              a) —CH<sub>2</sub>—CH<sub>3</sub>, or
b) —CH<sub>2</sub>-cyclopropyl;
         -CH<sub>2</sub>-SO<sub>2</sub>-phenyl substituted by zero (0) or one
   (1) R<sub>11</sub>, or
d) —CH<sub>2</sub>—SO<sub>2</sub>—het;
                                                                                           wherein Ro is
wherein het is a 5-, 6- or 7-membered saturated or
                                                                                               a) -NR_{12}SO_2—het,
   unsaturated ring containing from one (1) to three (3)
                                                                                              b) -NR<sub>12</sub>SO<sub>2</sub>-phenyl substituted by zero (0) or one (1)
   heteroatoms selected from the group consisting of
   nitrogen, oxygen and sulfur; and including any bicyclic 40
                                                                                              c) -CH<sub>2</sub>-SO<sub>2</sub>-phenyl substituted by zero (0) or one
   group in which any of the above heterocyclic rings is
                                                                                                  (1) R<sub>11</sub>, or
   fused to a benzene ring or another heterocycle; substi-
                                                                                              d) -CH<sub>2</sub>-SO<sub>2</sub>-het;
   tuted by zero (0) or one (1) R<sub>10</sub>;
                                                                                           wherein het is a 5-, 6- or 7-membered saturated or
wherein R<sub>10</sub> is
                                                                                              unsaturated ring containing from one (1) to three (3)
                                                                                  45
   a) —CH<sub>3</sub>,
b) —CN,
                                                                                              heteroatoms selected from the group consisting of
                                                                                              nitrogen, oxygen and sulfur; and including any bicyclic
   c) -OH,
                                                                                              group in which any of the above heterocyclic rings is
   d) —C(0)OC_2H_5,
                                                                                              fused to a benzene ring or another heterocycle; substi-
   e) —CF<sub>3</sub>,
                                                                                              tuted by zero (0) or one (1) R<sub>10</sub>;
   f) - NH_2, or
                                                                                  50
                                                                                           wherein R<sub>10</sub> is
   g) ---C(O)---NH<sub>2</sub>;
                                                                                              a) —CH<sub>3</sub>,
b) —CN,
c) —OH, or
wherein R<sub>11</sub> is
   a) ---CN,
   b) —F,
c) —OH, or
                                                                                              d) -C(O)OC_2H_5;
                                                                                          wherein R<sub>11</sub> is
   d) —NO_2;
                                                                                              a) —CN,
b) —F,
wherein R<sub>12</sub> is
   a) —H, or
                                                                                              c) -OH, or
   b) —CH<sub>3</sub>;
                                                                                              d) -NO<sub>2</sub>;
or a pharmaceutically acceptable salt thereof.
                                                                                          wherein R<sub>12</sub> is
The present invention more particularly provides:
                                                                                              a) ---H, or
A compound of the formula I
                                                                                              b) ---CH<sub>3</sub>;
wherein R<sub>1</sub> is H-;
                                                                                          or a pharmaceutically acceptable salt thereof.
wherein R2 is
                                                                                          The present invention provides for such compounds
   a) C<sub>3</sub>-C<sub>5</sub> alkyl,
                                                                                       wherein het is the following, substituted by zero (0) or one
   b) phenyl-(CH<sub>2</sub>)<sub>2</sub>—, or
                                                                                       (1) R_{10},
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a) 2-pyridinyl,
                                                                                                 f) tetrahydropyran-3-yl, or
                                                                                                 g) benzimidazol-2-yl;
   b) imidazol-2-yl,
                                                                                              wherein R<sub>10</sub> is
   c) imidazol-4-yl,
                                                                                                 a) —CH<sub>3</sub>;
   d) benzimidazol-2-yl,
                                                                                             wherein R<sub>11</sub> is
   e) quinolin-8-yl,
                                                                                                 a) —CN,
   f) quinolin-2-yl,
                                                                                                 b) --- F or
                                                                                                 c) -NO<sub>2</sub>;
   g) pyrimidin-2-yl,
                                                                                             or a pharmaceutically acceptable salt thereof.
   h) quinazolin-2-yl,
                                                                                             Most particularly, the present invention provides for the
   i) purin-6-yl,
                                                                                          compound of the formula VI
   j) thiazol-2-yl,
                                                                                             wherein R2 is
                                                                                                 a) H<sub>3</sub>C—(CH<sub>2</sub>)<sub>2</sub>—
   k) thiazol-4-yl,
                                                                                                 b) phenyl-(CH<sub>2</sub>)<sub>2</sub>—,
   l) 2-pyrazolyl,
                                                                                     15
                                                                                                 c) (CH_3)_2CH—CH_2—, or
   m) 2-pyrazinyl,
                                                                                                 d) pentyl;
   n) tetrahydropyran-4-yl, or
                                                                                             wherein R<sub>3</sub> is the moiety of formula X
   o) tetrahydropyran-3-yl.
                                                                                             wherein R6 is
   Also more particularly, the present invention provides for
                                                                                                 a) H<sub>3</sub>C--(CH<sub>2</sub>)<sub>2</sub>--,
the compound of the formula I
                                                                                                 b) phenyl-(CH<sub>2</sub>)<sub>2</sub>—,
                                                                                                 c) (CH_3)_2CH—CH_2—, or
   wherein R<sub>1</sub> is H—;
                                                                                                 d) pentyl;
   wherein R2 is
                                                                                             wherein R7 is
      a) H<sub>3</sub>C—(CH<sub>2</sub>)<sub>2</sub>—,
      b) phenyl-(CH<sub>2</sub>)<sub>2</sub>-
                                                                                                 a) CH<sub>3</sub>---CH<sub>2</sub>---, or
                                                                                     25
                                                                                                 b) cyclopropyl;
      c) (CH<sub>3</sub>)<sub>2</sub>CH—CH<sub>2</sub>, or
      d) pentyl;
                                                                                             wherein R<sub>9</sub> is
                                                                                                 a) —NHSO<sub>2</sub>-phenyl substituted by one (1) R_{11}, or b) —NHSO<sub>2</sub>—het;
   or wherein R<sub>1</sub> and R<sub>2</sub> taken together are a double bond;
   wherein R<sub>3</sub> is the moiety of formula X
                                                                                             wherein het is the following, substituted by zero (0) or one
   wherein R<sub>4</sub> is
                                                                                                 (1) R_{10}
      a) phenyl,
                                                                                                 a) imidazol-4-yl, or
      b) het,
                                                                                                 b) quinolin-8-yl;
      c) cyclopropyl,
                                                                                             wherein R<sub>10</sub> is ---CH<sub>3</sub>;
      d) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—,
e) het—SO<sub>2</sub>NH—,
                                                                                             wherein R<sub>11</sub> is a) —CN, or b) —F.
                                                                                     35
      f) Br—,
                                                                                             Also, most particularly, the present invention provides for
      h) HO<sub>3</sub>S(CH<sub>2</sub>)<sub>2</sub>—N(CH<sub>3</sub>)—C(O)—(CH<sub>2</sub>)<sub>6</sub>—C(O)—
                                                                                         the compound of the formula VII
         NH-;
                                                                                             wherein R<sub>3</sub> is the moiety of formula X
   wherein R<sub>5</sub> is —H;
                                                                                             wherein R4 is
   wherein R<sub>6</sub> is
                                                                                                 a) phenyl,
     a) R<sub>4</sub>—(CH<sub>2</sub>)<sub>n</sub>—CH(R<sub>8</sub>)—,
b) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—CH<sub>2</sub>—,
c) H<sub>3</sub>C—(CH<sub>2</sub>)<sub>2</sub>—,
                                                                                                b) het,
                                                                                                 c) cyclopropyl,
                                                                                     45
                                                                                                d) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—,
e) het—SO<sub>2</sub>NH—,
      d) phenyl-(CH_2)_2—,
      e) (CH_3)_2CH—CH_2—, or
                                                                                                f) Br—,
g) N<sub>3</sub>—, or
      f) pentyl;
   wherein n is zero (0), one (1) or two (2);
                                                                                                 h) HO_3S(CH_2)_2—N(CH_3)—C(O)—(CH_2)_6—C(O)—
   wherein R7 is
                                                                                                   NH-:
                                                                                             wherein R<sub>6</sub> is
      a) cyclopropyl, or
      b) CH<sub>3</sub>—CH<sub>2</sub>—;
                                                                                                a) R<sub>4</sub>—(CH<sub>2</sub>)<sub>n</sub>—CH(R<sub>8</sub>)—, or
b) H<sub>3</sub>C—[O(CH<sub>2</sub>)<sub>2</sub>]<sub>2</sub>—CH<sub>2</sub>—;
  wherein R<sub>8</sub> is
      a) -CH_2-CH_3, or
                                                                                             wherein R<sub>7</sub> is cyclopropyl;
      b) —CH<sub>2</sub>-cyclopropyl;
                                                                                             wherein R<sub>8</sub> is
                                                                                                a) CH<sub>2</sub>---CH<sub>3</sub>, or
  wherein Ro is
      a) —NHSO<sub>2</sub>—het, or
                                                                                                b) —CH<sub>2</sub>-cyclopropyl;
            -NHSO<sub>2</sub>-phenyl substituted by zero (0) or one (1)
                                                                                             wherein Ro is
                                                                                                 a) -NHSO<sub>2</sub>-het, or
                                                                                                b) -NHSO<sub>2</sub>-phenyl substituted by one (1) R<sub>11</sub>;
  wherein het is the following, substituted by zero (0) or one
                                                                                             wherein n is zero (0), one (1) or two (2);
      (1) R_{10}
     a) 2-pyridinyl,
                                                                                             wherein het is the following, substituted by zero (0) or one
     b) imidazol-2-yl,
                                                                                                (1) R<sub>10</sub>,
     c) imidazol-4-yl,
                                                                                    65
                                                                                                a) imidazol-4-yl,
     d) quinolin-8-yl,
                                                                                                b) imidazol-2-yl,
     e) tetrahydropyran-4-yl,
                                                                                                c) quinolin-8-yl,
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d) tetrahydropyran-3-yl,
                                                                                   c) imidazol-4-yl,
      e) tetrahydropyran-4-yl,
                                                                                   d) benzimidazol-2-yl,
      f) 2-pyridinyl, or
                                                                                   e) quinolin-8-vl.
      g) benzimidazol-2-yl;
                                                                                  f) quinolin-2-yl,
   wherein R<sub>10</sub> is ---CH<sub>3</sub>;
                                                                                   g) pyrimidin-2-yl,
   wherein R<sub>11</sub> is
                                                                                  h) quinazolin-2-yl,
      a) -NO<sub>2</sub>,
                                                                                  i) purin-6-yl,
      b) ---F, or
      c) -CN;
                                                                                  j) thiazol-2-yl,
   or a pharmaceutically acceptable salt thereof.
                                                                                  k) thiazol-4-yl,
   The present invention also provides:
                                                                                  1) 2-pyrazolyl,
   A compound of the formula II
                                                                                  m) 2-pyrazinyl,
   wherein R<sub>10</sub> and R<sub>20</sub> taken together are
                                                                                  n) tetrahydropyran-4-yl, or
      a) the moiety of formula III, or
                                                                                  o) tetrahydropyran-3-yl.
      b) the moiety of formula IV;
                                                                                  More particularly, the present invention provides for the
   wherein p is four (4);
                                                                               compound of the formula II
   wherein R_1 is —H;
                                                                                  wherein R<sub>10</sub> and R<sub>20</sub> taken together are
   wherein R2 is
                                                                                      a) the moiety of formula III, or
      a) H-
                                                                                      b) the moiety of formula IV;
      b) CH<sub>3</sub>O--, or
                                                                                  wherein p is four (4);
      c) CH<sub>3</sub>O--[(CH<sub>2</sub>)<sub>2</sub>O]<sub>3</sub>--;
                                                                                  wherein R_1 is —H;
   wherein R<sub>3</sub> is the moiety of formula V
                                                                                  wherein R2 is
   wherein R4 is
                                                                           25
                                                                                      a) CH<sub>3</sub>O---, or
      a) cyclopropyl, or
                                                                                      b) CH<sub>3</sub>O[(CH<sub>2</sub>)<sub>2</sub>O]<sub>3</sub>—;
      b) —CH<sub>2</sub>—CH(CH<sub>3</sub>)<sub>2</sub>;
                                                                                  wherein R<sub>3</sub> is the moiety of formula V
   wherein R<sub>5</sub> is
                                                                                  wherein R4 is
      a) -NR<sub>9</sub>SO<sub>2</sub>-phenyl substituted by zero (0) or one (1)
                                                                                      a) cyclopropyl, or
                                                                                     b) -CH_2-CH(CH_3)_2;
      b) -NR<sub>9</sub>SO<sub>2</sub>-het,
c) -CH<sub>2</sub>-SO<sub>2</sub>-phenyl substituted by zero (0) or one
                                                                                   wherein R<sub>5</sub> is
                                                                                      a) -NR<sub>9</sub>SO<sub>2</sub>-phenyl substituted by zero (0) or one (1)
         (1) R_6, or
                                                                                     R<sub>6</sub>,
b) —NR<sub>9</sub>SO<sub>2</sub>—het,
c) —CH<sub>2</sub>—SO<sub>2</sub>-phenyl substituted by zero (0) or one
      d) -CH<sub>2</sub>-SO<sub>2</sub>-het;
   wherein R6 is
                                                                           35
      a) ---CN,
                                                                                        (1) R<sub>6</sub>, or
      b) ---F,
                                                                                      d) —CH<sub>2</sub>—SO<sub>2</sub>—het;
      c) —CH<sub>3</sub>,
                                                                                  wherein R<sub>6</sub> is
      d) —COOH, or
                                                                                     a) —CN,
b) —F,
      e) --OH;
   wherein het is a 5-, 6- or 7-membered saturated or
                                                                                     c) ---CH<sub>3</sub>, or
      unsaturated ring containing from one (1) to three (3)
                                                                                     d) —COOH;
      heteroatoms selected from the group consisting of
                                                                                  wherein het is the following, substituted by zero (0) or one
      nitrogen, oxygen and sulfur; and including any bicyclic
                                                                                     (1) R_{7},
      group in which any of the above heterocyclic rings is 45
                                                                                     a) imidazol-4-yl,
      fused to a benzene ring or another heterocycle; substi-
                                                                                     b) quinolin-8-yl,
      tuted by zero (0), one (1) or two (2) R_7;
                                                                                     c) 2-pyridinyl, or
   wherein R7 is
                                                                                     d) 4-pyridinyl;
     a) —CH<sub>3</sub>,
b) —CN,
                                                                                  wherein R<sub>7</sub> is —CH<sub>3</sub>;
                                                                                  wherein R<sub>8</sub> is
      c) —C(0)OC_2H_5, or
      d) —OH;
                                                                                     a) —H, or
                                                                                     b) —(CH<sub>2</sub>)<sub>2</sub>—CH<sub>3</sub>;
   wherein R<sub>8</sub> is
      a) ---H,
                                                                                  wherein R<sub>9</sub> is
                                                                                     a) —H, or
b) —CH<sub>3</sub>;
      b) -(CH<sub>2</sub>)<sub>2</sub>--CH<sub>3</sub>,
      c) -CH2-cyclopropyl, or
      d) --CH<sub>2</sub>-phenyl;
                                                                                  or a pharmaceutically acceptable salt thereof.
   wherein Ro is
                                                                                  Most particularly, the present invention provides for the
                                                                           compound of the formula VIII
     a) —H, or
     b) ---CH<sub>3</sub>;
                                                                                  wherein R<sub>3</sub> is the moiety of formula V
  or a pharmaceutically acceptable salt thereof.
                                                                                  wherein R4 is
  The present invention provides for such compounds
                                                                                     a) cyclopropyl,
wherein het is the following, substituted by zero (0) or one
                                                                                     b) -CH_2-CH(CH_3)_2;
(1) R<sub>2</sub>,
                                                                                  wherein R<sub>5</sub> is
   a) 2-pyridinyl,
                                                                                     a) —NR<sub>9</sub>SO<sub>2</sub>-phenyl substituted by zero (0) or one (1)
  b) imidazol-2-yl,
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b) -NR<sub>9</sub>SO<sub>2</sub>-bet, or
                                                                                        b) 2-pyridinyl, or
      c) -CH<sub>2</sub>-SO<sub>2</sub>-phenyl;
                                                                                        c) quinolin-8-yl;
                                                                                     wherein R<sub>10</sub> is,
   wherein R<sub>6</sub> is
                                                                                       a) —CH<sub>3</sub>,
      a) ---CN, or
      b) —F;
                                                                                        b) —CN,
                                                                                       c) --- CF<sub>3</sub>,
   wherein het is the following, substituted by zero (0) or one
                                                                                        d) -NH2 or
      (1) R_{7},
                                                                                        e) ---C(O)---NH<sub>2</sub>;
      a) 2-pyridinyl,
      b) 4-pyridinyl, or
                                                                                     wherein R<sub>11</sub> is CN.
      c) imidazol-4-yl;
                                                                                    The present invention also provides:
   wherein R<sub>7</sub> is —CH<sub>3</sub>;
                                                                                    A compound of the formula XI
   wherein R<sub>8</sub> is
                                                                                    wherein R_1 is -(CH_2)_p--CH(R_2)-(CH_2)_o-Ar_1;
      a) —H, or
b) —(CH<sub>2</sub>)<sub>2</sub>—CH<sub>3</sub>;
                                                                                    wherein R2 is
                                                                                       a) -C_1-C_5 alkyl, or
b) -(CH_2)_q-cycloalkyl;
                                                                             15
   wherein R9 is
      a) -H, or
                                                                                    wherein Ar<sub>1</sub> is
      b) ---CH3;
                                                                                        a) phenyl substituted by zero (0) or one (1) R<sub>3</sub>, or
   or a pharmaceutically acceptable salt thereof.
                                                                                        b) phenyl substituted by -meta-NHSO<sub>2</sub>Ar<sub>2</sub>;
   Also, most particularly, the present invention provides for 20
                                                                                    wherein Ar, is
the compound of the formula IX
                                                                                        a) phenyl substituted by zero (0) or one (1) R<sub>3</sub>, or
   wherein R<sub>1</sub> is H-:
                                                                                        b) het;
   wherein R<sub>2</sub> is
                                                                                    wherein het is a 5-, 6- or 7-membered saturated or
      a) CH_3O—, or
                                                                                        unsaturated ring containing from one (1) to three (3)
                                                                             25
      b) CH<sub>3</sub>O—[(CH<sub>2</sub>)<sub>2</sub>O]<sub>3</sub>—;
                                                                                        heteroatoms selected from the group consisting of
   wherein R<sub>3</sub> is the moiety of formula V
                                                                                        nitrogen, oxygen and sulfur; and including any bicyclic
                                                                                        group in which any of the above heterocyclic rings is
   wherein R<sub>4</sub> is cyclopropyl;
                                                                                        fused to a benzene ring or another heterocycle; substi-
   wherein R<sub>5</sub> is -NHSO<sub>2</sub>-het;
                                                                                        tuted by zero (0) or one (1) R<sub>4</sub>;
   wherein het is the following, substituted by zero (0) or one 30
                                                                                    wherein R<sub>3</sub> is
      (1) R_{7},
                                                                                       a) —CN,
b) —F,
      a) imidazol-4-yl,
     b) 2-pyridinyl, or
                                                                                       c) -OH, or
      c) quinolin-8-yl;
                                                                                        d) —NO<sub>2</sub>;
   wherein R7 is -CH3.
                                                                             35
   The present invention also provides for the compound of
                                                                                    wherein R₄ is
the formula VI
                                                                                        a) —CH<sub>3</sub>,
                                                                                       b) —CN,
   wherein R2 is
                                                                                       c) —OH,
      a) H<sub>3</sub>C—CH<sub>2</sub>—,
b) H<sub>3</sub>C—(CH<sub>2</sub>)<sub>2</sub>-
                                                                                       d) -C(0)OC_2H_5,
                                                                             40
                                                                                       e) --- CF<sub>3</sub>, or
      c) cyclopropyl-(CH<sub>2</sub>)<sub>2</sub>—,
                                                                                       f) ---NH<sub>2</sub>;
      d) F-phenyl-(CH<sub>2</sub>)<sub>2</sub>-
      e) het-SO2NH-phenyl-,
                                                                                    wherein n is zero (0) to eight (8), inclusive;
     f) (H<sub>3</sub>C)<sub>2</sub>HC—CH<sub>2</sub>,
                                                                                    wherein o is zero (0) to three (3), inclusive;
     g) phenyl-(CH<sub>2</sub>)<sub>2</sub>--, or
                                                                                    wherein p is zero (0) to three (3), inclusive;
     h) F_3C-(CH_2)_2-
                                                                                    wherein q is zero (0) to three (3), inclusive; or
   wherein R<sub>3</sub> is the moiety of formula X
                                                                                    a pharmaceutically acceptable salt thereof.
   wherein R6 is
                                                                                    More particularly, the present invention provides:
      a) H<sub>2</sub>C—CH<sub>2</sub>-
                                                                                    The compound wherein R_1 is -CH(R_2)-Ar_1;
     b) H<sub>3</sub>C—(CH<sub>2</sub>)<sub>2</sub>-
                                                                                    wherein R2 is
     c) cyclopropyl-(CH<sub>2</sub>)<sub>2</sub>--,
                                                                                       a) --CH_2---CH_3, or
     d) F-phenyl-(CH<sub>2</sub>)<sub>2</sub>-
                                                                                       b) -t-butyl;
     e) het-SO<sub>2</sub>NH-phenyl,
                                                                                    wherein Ar<sub>1</sub> is phenyl substituted by -meta-NHSO<sub>2</sub>Ar<sub>2</sub>;
     f) (H<sub>3</sub>C)<sub>2</sub>HC—CH<sub>2</sub>,
                                                                                    wherein Ar<sub>2</sub> is 2-pyridinyl substituted by one (1) R<sub>4</sub>;
     g) phenyl-(CH<sub>2</sub>)<sub>2</sub>-, or
                                                                                    wherein R<sub>4</sub> is a) —CN, or
     h) F_3C-(CH_2)_2-;
   wherein R7 is
     a) H<sub>3</sub>C--CH<sub>2</sub>---,
                                                                                       b) —CF<sub>3</sub>;
     b) t-butyl, or
                                                                                    wherein n is two (2) to four (4) inclusive.
     c) cyclopropyl
                                                                                    The present invention also provides:
                                                                                    A process for producing a compound of the formula W-10
   wherein R<sub>o</sub> is
     a) -NHSO<sub>2</sub>-het, or
                                                                                    wherein R, is
     b) -NHSO<sub>2</sub>-phenyl substituted by one (1) R<sub>11</sub>;
                                                                                       a) n-propyl, or
   wherein het is the following, substituted by zero (0) or one 65
                                                                                       b) phenethyl;
     (1) R<sub>10</sub>,
                                                                                    which comprises the steps of:
     a) imidazol-4-yl,
                                                                                       a) treating a compound of the formula W-9
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wherein X<sub>A</sub> is as defined above, with TiCl<sub>4</sub>;

- b) treating the product of step a) with an amine base;
   and
- c) reacting the product of step b) with 4-heptanone or propylphenethylketone to yield the compound of 5 formula W-10;

The process which further comprises the steps of:

d) treating the compound of formula W-10 with sodium hydride or potassium t-butoxide to obtain a compound of formula W-11

wherein R, is

- a) n-propyl, or
- b) phenethyl;
- e) hydrogenating the compound of formula W-11 to obtain the compound of formula W-12

wherein R<sub>1</sub> is as defined above;

- f) treating the compound of formula W-12 with a sulfonyl chloride of formula D-7
- wherein  $R_4$  is 5-trifluoromethyl-2-pyridinyl, in an organic solvent in the presence of an organic base to obtain a compound of the formula W-13

wherein R<sub>1</sub> is as defined above.

The present invention also provides:

A process for producing a compound of the formula X-10 25

wherein  $R_1$  is

- a) n-propyl, or
- b) phenethyl;

which comprises the steps of:

a) treating a compound of the formula X-9

wherein X<sub>A</sub> is as defined above, with TiCl<sub>4</sub>;

- b) treating the product of step a) with an amine base; and
- c) reacting the product of step b) with 4-heptanone or propylphenethylketone to yield the compound of formula 35 X-10;

The process which further comprises the steps of:

d) treating the compound of formula X-10 with sodium hydride or potassium t-butoxide to obtain a compound of formula X-11

wherein R<sub>1</sub> is

- a) n-propyl, or
- b) phenethyl;
- e) hydrogenating the compound of formula X-11 to obtain a compound of formula X-12 wherein R<sub>1</sub> is as defined above;
- f) treating the compound of formula X-12 with a sulfonyl chloride of formula D-7
- wherein R<sub>4</sub> is 5-trifluoromethyl-2-pyridinyl, in an organic solvent in the presence of an organic base to obtain a compound of the formula X-13

wherein  $R_1$  is as defined above.

The present invention also provides:

A process for producing a compound of the formula  $_{55}$  GGG-10

wherein R<sub>1</sub> is

- a) n-propyl, or
- b) phenethyl;

which comprises the steps of:

- a) treating a compound of the formula GGG-9 wherein X<sub>A</sub> is as defined above, with TiCl<sub>4</sub>;
  - b) treating the product of step a) with an amine base;
  - c) reacting the product of step b) with 4-heptanone or 65 1-phenyl-3-hexanone to yield the compound of formula GGG-10.

The process which further comprises the steps of:

d) treating the compound of formula GGG-10 with sodium hydride or potassium t-butoxide to obtain a compound of formula GGG-11

wherein R<sub>1</sub> is

- a) n-propyl, or
- b) phenethyl;
- e) hydrogenating the compound of formula GGG-11 to obtain a compound of formula GGG-12

wherein R<sub>1</sub> is as defined above;

f) treating the compound of formula GGG-12 with a sulfonyl chloride of formula D-7

wherein R4 is

- a) 5-trifluoromethyl-2-pyridinyl, or
- b) 5-cyano-2-pyridinyl, in an organic solvent in the presence of an organic base to obtain a compound of the formula GGG-13A

wherein  $R_1$  is as defined above.

A process for producing a compound of the formula HHH-10

wherein R<sub>1</sub> is

- a) n-propyl, or
- b) phenethyl;

which comprises the steps of:

a) treating a compound of the formula HHH-9

wherein X<sub>A</sub> is as defined above, with TiCl<sub>4</sub>;

- b) treating the product of step a) with an amine base;
   and
- c) reacting the product of step b) with 4-heptanone or 1-phenyl-3-hexanone to yield the compound of formula HHH-10.

The process which further comprises the steps of:

d) treating the compound of formula HHH-10 with sodium hydride or potassium t-butoxide to obtain a compound of formula HHH-11

wherein R<sub>1</sub> is

- a) n-propyl, or
- b) phenethyl;
- e) hydrogenating the compound of formula HHH-11 to obtain a compound of formula HHH-12

wherein R<sub>1</sub> is as defined above;

f) treating the compound of formula HHH-12 with a sulfonyl chloride of formula D-7

wherein R<sub>4</sub> is

- a) 5-trifluoromethyl-2-pyridinyl, or
- b) 5-cyano-2-pyridinyl, in an organic solvent in the presence of an organic base to obtain a compound of the formula HHH-13A

wherein R<sub>1</sub> is as defined above.

A process for producing a compound of the formula III-10 wherein  $R_1$  is

- a) n-propyl, or
- b) phenethyl;

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which comprises the steps of:

a) treating a compound of the formula III-9

wherein X<sub>A</sub> is as defined above, with TiCl<sub>4</sub>;

- b) treating the product of step a) with an amine base; and
- c) reacting the product of step b) with 4-heptanone or 1-phenyl-3-hexanone to yield the compound of formula III-10.

The process which further comprises the steps of:

d) treating the compound of formula III-10 with sodium hydride or potassium t-butoxide to obtain a compound of formula III-11

- a) n-propyl, or
- b) phenethyl;

wherein R<sub>1</sub> is

e) hydrogenating the compound of formula III-11 to obtain a compound of formula III-12

wherein R<sub>1</sub> is as defined above;

f) treating the compound of formula III-12 with a sulfonyl chloride of formula D-7

wherein R<sub>4</sub> is

a) 5-trifluoromethyl-2-pyridinyl, or

b) 5-cyano-2-pyridinyl, in an organic solvent in the presence of an organic base to obtain a compound of the formula III-13A

wherein R<sub>1</sub> is as defined above.

A process for producing a compound of the formula

wherein R, is

- a) n-propyl, or
- b) phenethyl;

which comprises the steps of:

a) treating a compound of the formula JJJ-9

wherein X<sub>A</sub> is as defined above, with TiCl<sub>4</sub>;

- b) treating the product of step a) with an amine base;
- c) reacting the product of step b) with 4-heptanone or 1-phenyl-3-hexanone to yield the compound of formula JJJ-10.

The process which further comprises the steps of:

d) treating the compound of formula JJJ-10 with 30 fluorine, chlorine, bromine, and iodine sodium hydride or potassium t-butoxide to obtain a compound of formula JJJ-11

wherein R<sub>1</sub> is

- a) n-propyl, or
- b) phenethyl;
- e) hydrogenating the compound of formula JJJ-11 to obtain a compound of formula JJJ-12

wherein R<sub>1</sub> is as defined above;

sulfonyl chloride of formula D-7

wherein R<sub>4</sub> is

- a) 5-trifluoromethyl-2-pyridinyl, or
- b) 5-cyano-2-pyridinyl, in an organic solvent in the presence of an organic base to obtain a compound of 45 the formula JJJ-13A

wherein R<sub>1</sub> is as defined above.

The present invention most preferably provides:

The compound of formula VI wherein R<sub>2</sub> is

- a)  $H_3C-(CH_2)_2$ , or
- b) phenyl-(CH<sub>2</sub>)<sub>2</sub>—;

wherein R<sub>3</sub> is the moiety of formula X;

wherein R<sub>6</sub> is

- a)  $H_3C$ — $(CH_2)_2$ —, or
- b) phenyl- $(CH_2)_2$ —;

wherein R7 is

- a)  $H_3C CH_2$ —, or
- b) t-butyl;

wherein R<sub>o</sub> is —NHSO<sub>2</sub>—het;

wherein het is the following, substituted by one (1) R<sub>10</sub>,

- a) imidazol-4-yl, or
- b) 2-pyridinyl;

wherein R<sub>10</sub> is,

- a) —CH<sub>3</sub>, b) —CN, or

c) --- CF<sub>3</sub>.

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The compounds of the present invention are named according to the IUPAC or CAS nomenclature system.

The carbon atoms content of various hydrocarboncontaining moieties is indicated by a prefix designating the minimum and maximum number of carbon atoms in the moiety, i.e., the prefix  $C_i$ — $C_j$  indicates a moiety of the integer "i" to the integer "j" carbon atoms, inclusive. Thus, for example, C<sub>1</sub>-C<sub>3</sub> alkyl refers to alkyl of one to three carbon atoms, inclusive, or methyl, ethyl, propyl, and isopropyl, 10 straight and branched forms thereof.

Also, the carbon atom content of various hydrocarboncontaining moieties of the present invention is indicated by a subscripted integer representing the number of carbon and hydrogen atoms in the moiety, e.g., "C<sub>n</sub>H<sub>2n</sub>" indicates a 15 moiety of the integer "n" carbon atoms, inclusive, and the integer "2n" hydrogen atoms, inclusive. Thus, for example, "C<sub>n</sub>H<sub>2n</sub>" wherein n is one to three carbon atoms, inclusive, and two to six hydrogen atoms, inclusive, or methyl, ethyl, propyl and isopropyl, and all isomeric, straight and branched 20 forms thereof.

Examples of alkyl of one to nine carbon atoms, inclusive, are methyl, ethyl, propyl, butyl, pentyl, hexyl, heptyl, octyl, and nonyl, and all isomeric forms thereof and straight and branched forms thereof.

Examples of alkenyl of one to five carbon atoms, inclusive, are ethenyl, propenyl, butenyl, pentenyl, all isomeric forms thereof, and straight and branched forms thereof.

By "halo" is meant the typical halogen atoms, such as

The compounds of formula I and II of the present invention inhibit retroviral proteinases and thus inhibit the replication of the virus. They are useful for treating patients infected with human immunodeficiency virus (HIV) which 35 results in acquired immunodeficiency syndrome (AIDS) and related diseases.

More particularly, the compounds of the present invention are useful as novel human retroviral protease inhibitors. Therefore, the compounds inhibit retroviral proteases and f) treating the compound of formula JJJ-12 with a 40 thus inhibit the replication of the virus. They are useful for treating human patients infected with a human retrovirus, such as human immunodeficiency virus (strains of HIV-1 or HIV-2) or human T-cell leukemia viruses (HTLV-I or HTLV-II) which results in acquired immunodeficiency syndrome (AIDS) and/or related diseases.

The capsid and replicative enzymes (i.e. protease, reverse transcriptase, integrase) of retroviruses are translated from the viral gag and pol genes as polyproteins that are further processed by the viral protease (PR) to the mature proteins 50 found in the viral capsid and necessary for viral functions and replication. If the PR is absent or nonfunctional, the virus cannot replicate. The retroviral PR, such as HIV-1 PR, has been found to be an aspartic protease with active site characteristics similar to those exhibited by the more com-55 plex aspartic protease, renin.

The term human retrovirus (HRV) includes human immunodeficiency virus type I, human immunodeficiency virus type II, or strains thereof, as well as human T cell leukemia virus 1 and 2 (HTLV-1 and HTLV-2) or strains apparent to 60 one skilled in the art, which belong to the same or related viral families and which create similar physiological effects in humans as various human retroviruses.

Patients to be treated would be those individuals: 1) infected with one or more strains of a human retrovirus as determined by the presence of either measurable viral antibody or antigen in the serum and 2) in the case of HIV, having either an asymptomatic HIV infection or a symp-

tomatic AIDS defining infection such as i) disseminated histoplasmosis, ii) isopsoriasis, iii) bronchial and pulmonary candidiasis including pneumocystic pneumonia iv) non-Hodgkin's lymphoma or v) Kaposi's sarcoma and being less than sixty years old; or having an absolute CD4+ lymphocyte count of less than 500/mm<sup>3</sup> in the peripheral blood. Treatment would consist of maintaining an inhibitory level of the compound used according to this invention in the patient at all times and would continue until the occurrence of a second symptomatic AIDS defining infection indicates 10 alternate therapy is needed.

More specifically, an example of one such human retrovirus is the human immunodeficiency virus (HIV, also known as HTLV-III or LAV) which has been recognized as the causative agent in human acquired immunodeficiency 15 syndrome (AIDS), P. Duesberg, Proc. Natl. Acad. Sci. USA, 86:755 (1989). HIV contains a retro viral encoded protease, HIV-I protease, that cleaves the fusion polypeptides into the functional proteins of the mature viral particle, E. P. Lillehoj, et al., J. Virology, 62:3053 (1988); C. Debuck, et al., Proc. 20 Natl. Acad. Sci., 84:8903 (1987). This enzyme, HIV-I protease, has been classified as an aspartyl protease and has a demonstrated homology to other aspartyl proteases such as renin, L. H. Pearl, et al., Nature 329:351 (1987); I. Katoh, et al., Nature 329:654 (1987). Inhibition of HIV-I protease 25 blocks the replication of HIV and thus is useful in the treatment of human AIDS, E. D. Clerq, J. Med. Chem. 29:1561 (1986). Inhibitors of HIV-I protease are useful in the treatment of HIV-infected individuals who are asymptomatic or symptomatic of AIDS.

Pepstatin A, a general inhibitor of aspartyl proteases, has been disclosed as an inhibitor of HIV-I protease, S. Seelmeier, et al., Proc. Natl. Acad. Sci. USA, 85:6612 (1986). Other substrate derived inhibitors containing reduced bond isosteres or statine at the scissle position have 35 also been disclosed, M. L. Moore, et al., Biochem. Biophys, Res. Commun. 159:420 (1989); S. Billich, et al., J. Biol. Chem. 263:17905 (1988); Sandoz, D. E. 3812-576-A.

Thus, the compounds of the present invention are useful acquired immunodeficiency disease syndrome (AIDS).

The compounds are also useful for treating non-human animals infected with a retrovirus, such as cats infected with feline leukemia virus. Other viruses that infect cats include, for example, feline infectious peritonitis virus, calicivirus, 45 rabies virus, feline immunodeficiency virus, feline parvovirus (panleukopenia virus), and feline chlamydia. Exact dosages, forms and modes of administration of the compounds of the present invention to non-human animals would be apparent to one of ordinary skill in the art, such as 50 a veterinarian.

The compounds of formula I and II of the present invention are prepared as described in the Charts, Preparations and Examples below, or are prepared by methods analogous thereto, which are readily known and available to one of 55 ordinary skill in the art of organic synthesis.

#### CHART A

Nitration of the cyclopropylphenyl ketone of formula A-1, which is commercially available, with fuming nitric acid at 60 -40° C. produces a ca. 2:1 mixture of isomers. The desired m-nitro compound of formula A-2 is easily separated from the crude mixture by recrystallization from methanol. Catalytic hydrogenation of the cyclopropyl-(3-nitrophenyl) methanone of formula A-2 with 10% platinum on carbon in methanol gives the aniline of formula A-3. The aniline is then coupled with benzenesulfonyl chloride using pyridine

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in methylene chloride to give the sulfonamide derivative of formula A-4. Reduction of the ketone with sodium borohydride in tetrahydrofuran and ethanol then produces the carbinol of formula A-5.

The dianion of the cyclooctylpyranone of formula A-6, prepared as described in Chart B, is formed using lithium diisopropyl amide in tetrahydrofuran at 0° C., and then alkylated with iodopropane to give the 10-propylcyclooctylpyranone of formula A-7. The cyclooctylpyranone of formula A-7 and the carbinol of the formula A-5 are then coupled using p-toluenesulfonic acid in methylene chloride to give the sulfonamide derivative of formula A-8.

#### CHART B

The commercially available amine of the formula B-1 is protected using benzyl chloroformate and sodium bicarbonate in THF/water solution to give the compound of formula B-2. The aldehyde of formula B-2 is then reacted with a Grignard reagent to give the secondary alcohol of formula B-3, wherein, e.g., R<sub>1</sub> is isobutyl. The known cyclooctylpyranone of formula B-4 is prepared by acylation of the trimethylsilyl enol ether of cyclooctanone with malonyl dichloride as described in R. Effenberger, T. Ziegler, K.-H. Schonwalder, T. Kesmarszky, B. Bauer Chem. Ber. 119:3394-3404 (1986). The alcohol of formula B-3 is then used to alkylate the cyclooctylpyranone of formula B-4 in refluxing toluene and p-toluenesulfonic acid to obtain the compound of the formula B-5, wherein, e.g., R<sub>1</sub> is isobutyl. At this point, the enantiomers of formula B-5 are separated using a chiral HPLC column. The benzyloxy protecting group is then cleaved using 10% Pd/C in cyclohexene to give the amine of formula B-6, wherein, e.g., R<sub>1</sub> is isobutyl, which is reacted with aryl sulfonyl chlorides to give the compounds of the formula B-7, wherein, e.g., R<sub>1</sub> is isobutyl and R<sub>2</sub> is 1-methylimidazole.

#### CHART C

3-Bromobenzyl alcohol of formula C-1, which is comfor treating diseases caused by retroviruses, such as human 40 mercially available, in tetrahydrofuran is treated with methyllithium, n-butyllithium and cyclopropanecarboxaldehyde in sequence at -78° C. The resulting solution is gradually warmed to room temperature and then heated at reflux affording the alcohol of formula C-2. The resulting alcohol, in dichloromethane, in the presence of molecular sieves, is treated with 4-hydroxy-5,6,7,8,9,10hexahydrocycloocta[b]pyran-2-one of formula C-8, prepared as described in Chart B, and p-toluenesulfonic acid. The solution is heated at reflux to afford the alcohol of formula C-3. The benzyl alcohol is treated with carbon tetrabromide and triphenylphosphine in dichloromethane at 0° C. to afford compounds of formula C-4 and C-5 as an inseparable mixture after an aqueous brine workup. The mixture is then treated with any thiol (e.g., thiophenol) and an organic base and heated at reflux to afford sulfides of the formula C-6. Finally treatment of the compounds of the formula C-6 with oxone in a mixture of tetrahydrofuran, methanol and water gives sulfones of formula C-7.

#### CHART D

This chart describes a generic procedure for the preparation of C-3a branched 5,6-dihydropyrones via aluminum chloride (AlCl<sub>3</sub>) mediated condensation with 3-nitrobenzaldehyde. Thus, the AlCl<sub>3</sub> catalyzed reaction of the compound of formula D-1, prepared as described below in the Preparations, (e.g., wherein R<sub>1</sub> is phenethyl or propyl; R<sub>2</sub> is phenethyl or propyl) with 3-nitrobenzaldehyde

(formula D-2), which is commercially available, provides compounds of formula D-3 (e.g., wherein R, is phenethyl or propyl; R2 is phenethyl or propyl). Subsequent reaction with trialkyl aluminums or Grignard reagents in the presence of cuprous bromide-dimethylsulfide complex (CuBr-Me<sub>2</sub>S) provides compounds of formula D-4 (e.g., wherein R<sub>1</sub> is phenethyl or propyl; R<sub>2</sub> is phenethyl or propyl; R<sub>3</sub> is ethyl or cyclopropyl). Transfer hydrogenation with Pd/C and ammonium formate provides compounds of formula D-5 propyl; R<sub>3</sub> is ethyl or cyclopropyl). Treatment of the compound of formula D-5 with sulfonyl chlorides of formula D-7, wherein R<sub>4</sub> is defined below, and pyridine in methylene chloride (CH<sub>2</sub>Cl<sub>2</sub>) provides compounds of formula D-6 (e.g., wherein R<sub>1</sub> is phenethyl or propyl; R<sub>2</sub> is phenethyl or 15 propyl; R<sub>3</sub> is ethyl or cyclopropyl; R<sub>4</sub> is 4-cyanophenyl, 4-fluorophenyl, 1-methylimidazol-4-yl, quinolin-8-yl, 2-pyridyl, 4-cyano-2-pyridyl, quinolin-2-yl, 2-hydroxyphenyl, 2-pyrimidyl, 2-quinazoline, 7H-purin-6yl, 1H-imidazol-2-yl, 1H-benzimidazol-2-yl or thiazol-2yl).

#### CHART E

Treatment of commercially available 4-hydroxy-6methyl-2-pyrone of formula E-1 with three equivalents of lithium diisopropylamide in tetrahydrofuran and hexamethylphosphoramide is followed by bromomethylcyclopropane to afford the compound of formula E-2. Reaction between the compound of formula E-2 and the compound of formula F-5, prepared as described in Chart F, in benzene with p-toluenesulfonic acid catalyst in the presence of molecular sieves affords the compound of formula E-3. Hydrogenolysis of the compound of formula E-3 in methanol with hydrogen and palladium on charcoal gives the free amine of formula E-4. Treatment of the compound of formula E-4 with two equivalents of pyridine in dichloromethane followed by one equivalent of 4-fluorobenzenesulfonyl chloride gives the compound of formula E-5 (wherein, e.g., R is 4-fluorophenyl) which is the compound: N-(3-{cyclopropyl-[6-(2-cyclopropyl-1-cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-fluorobenzenesulfonamide.

Under similar conditions, compounds of general formula E-5 are obtained by reacting amine E-4 with alkyl, aryl and heteroaryl sulfonyl chlorides in the presence of pyridine to give compounds of formula E-5 wherein R is alkyl, aryl or heteroaryl. Also, for example, the enantiomers of the compound of formula E-9 are separated chromatographically by chiral HPLC to give compounds of formula E-10 and E-11. Additional final compounds of the present invention of formula E-6, E-7, E-8, and E-12-E-16 are prepared using similar conditions.

## CHART F

Nitration of commercially available cyclopropyl phenyl ketone of formula F-1 with fuming nitric acid affords the 55 compound of formula F-2. Reduction of the compound of formula F-2 in methanol with hydrogen catalyzed by platinum on carbon gives the amine of formula F-3. The compound of formula F-3 is treated with benzylchloroformate and diisopropylethylamine in dichloromethane to give the 60 compound of formula F-4. Reduction of the compound of formula F-4 with sodium borohydride in tetrahydrofuran and ethanol gives the compound of formula F-5.

#### CHART G

The dianion of commercially available 4-hydroxy-6methyl-2-pyrone of formula G-0 is generated by deproto-

nation with two equivalents of lithium diisopropylamide in tetrahydrofuran and hexamethylphosphoramide. Alkylation with 2-(2-methoxy-ethoxy)-ethyl iodide, which is prepared from the commercially available alcohol by standard procedures, gives the compound of formula G-1. Reaction between the compound of formula G-1 and metabenzyloxycarbonylaminophenyl cyclopropyl carbinol, the compound of formula F-5, prepared as described in Chart F, in dichloromethane with p-toluenesulfonic acid catalyst in (e.g., wherein  $R_1$  is phenethyl or propyl;  $R_2$  is phenethyl or 10 the presence of molecular sieves gives the compound of formula G-2. Hydrogenolysis of the compound of formula G-2 in ethanol with hydrogen and palladium on charcoal gives the free amine of formula G-3. Treatment of the free amine of formula G-3 with two equivalents of pyridine in dichloromethane followed by one equivalent of 1-methylimidazole-4-sulfonyl chloride gives the compound of formula G-4, which is the compound: N-(3-{cyclopropyl-[4-hydroxy-6-(3-{2-methoxy-ethoxy}-propyl)-2-oxo-2Hpyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-20 sulfonamide.

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#### CHART H

Reaction between commercially available 4-hydroxy-6methyl-2-pyrone of formula H-0 and metabenzyloxycarbonylaminophenyl cyclopropyl carbinol, the title compound of formula F-5, prepared as described in Chart F, in dichloromethane with p-toluenesulfonic acid catalyst in the presence of molecular sieves gives the compound of formula H-1. Alkylation of trianion of the compound of formula H-1 generated from three equivalents of lithium diisopropylamide in tetrahydrofuran with ethyl bromide affords the compound of formula H-2. Treatment of the compound of formula H-2 with lithium diisopropylamide in tetrahydrofuran and 2-(2-methoxy-ethoxy)-ethyl iodide gives the compound of formula H-3. Hydrogenolysis of the compound of formula H-3 in ethanol with hydrogen and palladium on charcoal gives the free amine of formula H-4. Treatment of the free amine of formula H-4 with two equivalents of pyridine in dichloromethane followed by one equivalent of 1-methylimidazole-4-sulfonyl chloride gives the compound of formula H-5, which is the compound: N-(3-{cyclopropyl-[6-(1-ethyl-3-{2-methoxy-ethoxy}propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide. Under similar conditions, compounds of the present invention are obtained by reacting the amine of formula H-4 with alkyl, aryl and heteroaryl sulfonyl chlorides in the presence of pyridine to give additional sulfonamides of formula H-5.

#### CHART I

Treatment of the compound of formula H-2, prepared as described in Chart H, with three equivalents of lithium diisopropylamide in tetrahydrofuran and ethylene oxide gives the compound of formula I-1. Reaction of the compound of formula I-1 with triphenylphosphine and carbon tetrabromide in tetrahydrofuran gives the compound of formula I-2. Treatment of the compound of formula I-2 with sodium azide in aqueous ethanol gives the compound of formula I-3. Reaction of the compound of formula I-3 with hydrogen and palladium on charcoal in ethanol gives the compound of formula I-4. Treatment of the compound of formula I-4 with diisopropylethylamine in dichloromethane followed by 1-methylimidazole-4-sulfonyl chloride gives the compound of formula I-5. Reaction of the compound of formula I-5 with ammonia in methanol gives the compound of formula I-6, which is the compound: N-(3-{cyclopropyl-

[6-(1-ethyl-3-{1-methyl-1H-imidazole-4-sulfonylamino}propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl)-phenyl)-1-methyl-1H-imidazole-4-sulfonamide.

#### CHART J

Hydrogenolysis of the compound of formula I-1, prepared as described in Chart I, in ethanol with hydrogen and palladium on charcoal gives the compound of formula J-1. Treatment of the compound of formula J-1 with triphenylphosphine and carbon tetrabromide in tetrahydrofuran 10 gives the compound of formula J-2. Reaction of the compound of formula J-2 with pyridine in dichloromethane followed by 1-methylimidazole-4-sulfonyl chloride gives the compound of formula J-3, which is the compound: N-(3-{[6-(3-bromo-1-ethyl-propyl)-4-hydroxy-2-oxo-2Hpyran-3-yl]-cyclopropyl-methyl}-phenyl)-1-methyl-1Himidazole-4-sulfonamide. Treatment of the compound of formula J-3 with sodium azide in aqueous ethanol gives the compound of formula J-4, which is the compound: N-(3-{ ([6-(3-azido-1-ethyl-propyl)-4-hydroxy-2-oxo-2H-pyran-3-20 yl]-cyclopropyl-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide. Reaction of the compound of formula J-4 with hydrogen and palladium on charcoal in ethanol gives the compound of formula J-5. Treatment of the compound of formula J-5 with the triethylamine salt of suleptanic acid 25 (Anderson, B. D.; Conradi, R. A; Knuth, K. E.; J. Pharm. Sci. 74:365 (1985)) and 1,3-diisopropylcarbodiimide gives the compound of formula J-6, which is the compound:  $N-(3-\{cyclopropyl-[6-(1-ethyl-3-\{N-[8-(methyl-\{2-k-1])\})]\})$ sulfoethyl}-amino)-1,8-dioxooctyl]amino-propyl})-4- 30 hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide, sodium salt.

#### CHART K

The preparation of the compound of formula K-8, which is the compound: N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide is shown in Chart K. Reduction of commercially available tetrahydropyran-4-carboxylic acid of formula K-1 with borane in tetrahydrofuran provides the compound of formula K-2. The compound of formula K-2 is treated with p-toluenesulfonyl chloride to afford the corresponding tosylate of formula K-3, which is converted to the iodide of formula K-4 by treatment with potassium iodide in refluxing acetone. Alkylation of the dianion of commercially available 4-hydroxy-6-methyl-2-pyrone of formula K-10 with ethyl bromide in tetrahydrofuran and hexamethylphosphoric triamide gives the propyl derivative of formula K-9. The compound of formula K-4 is used to alkylate the compound of formula K-9 at the 6a position, giving the compound of formula K-5. The compound of formula K-5 is further alkylated at the 3 position, using carbinol of formula F-5, prepared as described in Chart F, giving the compound 55 of formula K-6. Removal of the benzyloxycarbonyl protecting group is accomplished using catalytic transfer hydrogenation, giving the amine of formula K-7. Treatment of the amine of formula K-7 with 1-methylimidazole-4compound of formula K-8.

#### CHART L

As shown in Chart L, the dianion of commercially available 4-hydroxy-6-methyl-2-pyrone of formula L-1 is gener- 65 ated by deprotonation with two equivalents of lithium diisopropylamide iп tetrahydrofuran

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hexamethylphosphoramide. Alkylation with benzyl bromide gives the compound of formula L-2, which is then treated with two equivalents of lithium diisopropylamide in tetrahydrofuran and hexamethylphosphoramide, followed by ethyl iodide to give the compound of formula L-3. Reaction between the compound of formula L-2 and the compound of formula F-5, prepared as described in Chart F, in benzene with p-toluenesulfonic acid catalyst in the presence of molecular sieves affords the compound of formula L-4, which is 3-[(3-benzyloxycarbonylaminophenyl)cyclopropyl-methyl]-6-(1-ethylphenethyl)-4-hydroxy-2Hpyran-2-one. Hydrogenolysis of the compound of formula L-4 in methanol using catalytic palladium on charcoal and ammonium formate or hydrogen gas gives the free amine of formula L-5, which is 3-[(3-aminophenyl)-cyclopropylmethyl]-6-(1-ethylphenethyl)-4-hydroxy-2H-pyran-2-one. Reacting the compound of formula L-5 and the appropriate sulfonyl chloride gives the final compounds of the present invention.

#### CHART M

As shown in Chart M, commercially available triethylene glycol monomethyl ether is treated with p-toluenesulfonyl chloride and pyridine to provide the tosylate of formula M-2, which is then used to alkylate commercially available 2,4dihydroxyacetophenone to give the compound of formula M-3. Condensation with diethyl carbonate yields the compound of formula M-4. Ring closure of the compound of formula M-4 to the compound of formula M-5 is accomplished by refluxing in acetic acid. The compound of formula M-5 is alkylated at the 3-position using the carbinol of formula F-5, prepared as described in Chart F, and catalytic p-toluenesulfonic acid to give the compound of formula M-6. Removal of the benzyloxycarbonyl protecting group is accomplished using catalytic transfer hydrogenation, giving the amine of formula M-7. Treatment of the amine with 1-methylimidazole-4-sulfonyl chloride in the presence of pyridine provides the final compound of formula M-8, which is N-(3-{Cyclopropyl-[7-(2-(2-methoxyethoxy)ethoxy)ethoxy)-4-hydroxycoumarin-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide.

#### CHART N

Nitration of cyclopropylphenyl ketone of formula N-1, which is commercially available, with fuming nitric acid at -40° C. produces a ca. 2:1 mixture of isomers. The desired meta-nitro compound of formula N-2 is easily separated from the crude mixture by recrystallization from methanol. Catalytic hydrogenation of cyclopropyl-(3-nitrophenyl) methanone of formula N-2 with 10% platinum on carbon in methanol at 0° C. provides the aniline of formula N-3. The product is isolated by filtration and concentration. The amino group is then protected using benzyl chloroformate and diisopropylethylamine in methylene chloride to give the ketone of formula N-4. The ketone is then reduced with sodium borohydride in 5:1 THF and ethanol to give the alcohol of formula N-5.

The compound of formula N-5 is then used to alkylate sulfonyl chloride in the presence of pyridine provides the 60 4-hydroxy-5,6,7,8,9,10-hexahydrocycloocta[b]pyran-2-one, which is prepared as described in R. Effenberger, T. Ziegler, K.-H. Schönzoalder, T. Kesmarsky, B. Bauer, Chem. Ber. 119:3394-3404 (1986), to give the compound of formula N-6. The preferred conditions for this alkylation reaction are p-toluene-sulfonic acid in refluxing methylene chloride with a Soxhlet extractor containing molecular sieves. Finally, the compound of formula N-7 is obtained by

cleaving the benzyl protective group in a transfer hydrogenation. Best results for this reactions are achieved with 10% Pd/C in neat cyclohexene.

#### CHART O

Treatment of the amine of formula O-1, prepared as described in Chart N, with sulfonyl chlorides and a base such as pyridine in dichloromethane gives the sulfonamides of formula O-2 wherein R<sub>60</sub> is, for example, 4-nitrophenyl. These sulfonamides are further modified by standard litera- 10 ture procedures as is apparent to those of ordinary skill in the art to give sulfonamides of formula O-3 wherein R<sub>61</sub> is, for example, 4-aminophenyl and other functional groups that are not readily available from readily available sulfonyl chlorides. For example, the nitro group of N-[3-[cyclopropyl 15 (5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b] pyran-3-yl)methyl]phenyl]-4-nitro-benzenesulfonamide is reduced by catalytic hydrogenation in ethyl acetate with palladium on carbon to give the amine in 4-amino-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H- 20 cycloocta[b]pyran-3-yl)methyl]phenyl]benzenesulfonamide. Also, the carboxylic acid of 3-[[[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2Hcycloocta[b]pyran-3-yl)methyl]phenyl]amino]sulfonyl]benzoic acid is esterified with methanol and catalytic 25 sulfuric acid to give the methyl ester in 3-[[[3-[cyclopropyl (5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b] pyran-3-yl)methyl]phenyl]amino]sulfonyl]-benzoic acid, methyl ester. Sulfonamides of formula O-3 are also obtained from compounds of formula O-2 by further elaboration of 30 reactive functional groups. For example, the amine of 3-amino-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]benzenesulfonamide is reacted with benzoyl chloride and a base such as pyridine to give the benzamide in N-[3-[[[3-35] [cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2Hcycloocta[b]pyran-3-yl)methyl]phenyl]amino]sulfonyl] henyl]-benzamide. Using commonly available sulfonyl chlorides, additional compounds of the present invention of formula II, wherein R<sub>10</sub> and R<sub>20</sub> is the moiety of formula IV, 40 are prepared.

The sulfonyl chlorides used to make the compounds of the present invention are readily prepared by methods described in the literature by those skilled in the art, as the following examples illustrate: Reaction of a suitable thiol with KHF<sub>2</sub> 45 in water/methanol with chlorine gas gives the sulfonyl fluoride (D. J. Brown, J. A. Hoskins, Aust. J. Chem. 25:2641 (1972)) which is then converted into the desired sulfonyl chloride (T. Norris, J. Chem. Soc., Perkin Trans. 1(11):1378 (Eng.) (1978)). Oxidation of a suitable thiol with chlorine in 50 water with ferric chloride (FeCl<sub>3</sub>) added gives the desired sulfonyl chloride (G. Pala, Ed. Sci. 13:461 (1958); W. J. Close, J. Amer. Chem. Soc. 82:1132 (1960)). Reaction of the heteroaromatic compound with fuming sulfuric acid gives a heteroaromatic sulfonic acid followed by treatment with 55 phosphorous-oxychloride (POCl<sub>2</sub>) and phosphorous chloride (PCl<sub>5</sub>) gives the desired sulfonyl chloride (V. Georgian, R. J. Harrison, L. L. Skaletzky, J. Org. Chem. 27:4571 (1962)). Reaction of a heteroaromatic compound with manganese dioxide (MnO<sub>2</sub>) and sodium sulfite (Na<sub>2</sub>SO<sub>3</sub>) in 60 water gives the desired sulfonic acid followed by treatment with POCl<sub>3</sub> and PCl<sub>5</sub> gives the desired sulfonyl chloride (N. A. Androva, Izvest. 455 (1972); J. O. Morley, J. Chem. Comm. 88 (1976)). Treatment of the appropriate heteroaromatic chloride with sodium sulfate and HCl in water gives 65 the desired sulfonic acid followed by treatment with POCl<sub>3</sub> and PCl<sub>5</sub> gives the desired sulfonyl chloride (T. R. Norton,

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J. Amer. Chem. Soc. 68:1330 (1946)). Treatment of the appropriate hydroxy compound with N, N-dimethylthiocarbonyl chloride (M. S. Newman, F. W. Hetzel, Org. Synth. Coll. Vol. IV:824 (1988); M. S. Newman, H. A. Karnes, J. Org. Chem. 31:3980 (1966)) followed by treatment of the resulting thiol, as described above, gives the desired sulfonyl chloride. Treatment of the appropriately protected thio-heteroaromatic compound with chlorine in acetic acid gives the desired sulfonyl chloride (Can. J. Chem. 55:421 (1977)). Using the literature procedures described above, the heteroaromatic sulfonyl chlorides of the present invention are prepared.

#### CHART P

The preferred procedure for the preparation of the heteroaryl sulfonamides of formula P-2 is described in Chart P. Sulfonation of the amine of formula P-1, prepared in Chart N, P-1 with various heteroarylsulfonyl chlorides of formula P-3 wherein R is, e.g., 2-pyridyl, 4-pyridyl, 5-cyanopyridin-2-yl, 2-pyrazinyl, 2-pyrimidinyl, 4,6-dimethylpyrimidin-2-yl, 4-methylpyrimidin-2-yl gives the sulfonamides of formula P-2 wherein R is the corresponding substituent.

#### CHART Q

Generated by sequential deprotonation with sodium hydride and n-butyl lithium in tetrahydrofuran at 0° C., the dianion of commercially available methyl acetoacetate is reacted with ketone of formula Q-1, prepared as described in Chart S (formula S-4). The resulting intermediate hydroxyester is cyclized with dilute aqueous hydroxide followed by aqueous hydrochloric acid to give the compound of formula Q-2. The compound of formula Q-2 is condensed with commercially available 3-nitrobenzaldehyde in tetrahydrofuran using aluminum trichloride as a catalyst followed by reaction of the intermediate benzylidene adduct with triethyl aluminum in the presence of copper bromide-dimethyl sulfide to provide the compound of formula Q-3. Catalytic transfer hydrogenation with Pd/C and ammonium formate in methanol affords the compound of formula Q-4. Treatment of the compound of formula Q-4 with the appropriate sulfonyl chloride and pyridine in dichloromethane provides the desired compound of formula Q-5 (wherein, e.g., R<sub>1</sub> is 5-cyano-2-pyridyl or 1-methylimidazol-4-yl).

#### CHART R

Catalytic hydrogenation of commercially available 3-nitropropiophenone of formula R-1 affords the amine of formula R-2. The amine of formula R-2 is treated with diisopropylethylamine and benzyl bromide to give the compound of formula R-3. The dianion of methyl acetoacetate, generated by treatment of commercially available methyl acetoacetate with sodium hydride and n-butyl lithium in tetrahydrofuran at 0° C., is reacted with the ketone of formula R-3. The intermediate hydroxy-ester is cyclized with dilute aqueous hydroxide followed by aqueous hydrochloric acid to give the compound of formula R-4. The compound of formula R-4 is condensed with 3-nitrobenzaldehyde in tetrahydrofuran using aluminum trichloride as a catalyst followed by reaction of the intermediate benzylidene adduct with triethyl aluminum in the presence of copper bromide-dimethyl sulfide to provide the compound of formula R-5. Catalytic hydrogenation with Pd/C affords the diamine of formula R-6. Treatment of the compound of formula R-6 with the appropriate sulfonyl chloride and pyridine in dichloromethane provides the desired compound of formula R-7 (wherein, e.g., R<sub>1</sub> is 5-cyano-2-pyridyl or 1-methylimidazol-4-yl).

Commercially available 4-pentenoic acid of formula S-1 is coupled with N,O-dimethylhydroxylamine using bis(2-oxo-3-oxazolidinyl)phosphinic chloride to afford the amide of formula S-2. The amide of formula S-2 is reacted with 3-butenyl magnesium bromide in tetrahydrofuran to give the ketone of formula S-3. The ketone of formula S-3 is treated with zinc metal, cuprous chloride and diiodomethane to provide the ketone of formula S-4 (also formula Q-1, see Chart Q above).

#### **CHART T**

The compound of formula T-2 (also formula D-1) (whose preparation is specifically described in Chart D and Prepa- 15 ration 17 above from commercially available methyl acetoacetate and 1-phenyl-3-hexanone (formula T-1)) is condensed with 3-nitrobenzaldehyde in tetrahydrofuran using aluminum trichloride as a catalyst followed by reaction of the intermediate benzylidene adduct with t-butylCu 20 (CN)ZnI, (the organometallic reagent derived from zinc metal, 2-iodo-2-methyl-propane, copper cyanide and lithium chloride) to provide the compound of formula T-3. (The preparation of the organometallic reagent is further described in the text corresponding to Preparation J above). 25 Catalytic transfer hydrogenation with Pd/C and ammonium formate in methanol affords the compound of formula T-4. Treatment of the compound of formula T-4 with the appropriate sulfonyl chloride and pyridine in dichloromethane provides the desired compound of formula T-5 (wherein, 30 e.g., R<sub>1</sub> is 5-cyano-2-pyridyl or 1-methylimidazol-4-yl).

#### CHART U

Commercially available 4-fluorohydrocinnamic acid of formula U-1 is coupled with N,O-dimethylhydroxylamine using diethyl cyanophosphonate to provide the amide of formula U-2. Treatment of the amide with n-propylmagnesium chloride yields the ketone of formula U-3. Condensation of the ketone with the dianion of methyl acetoacetate, followed by hydrolysis of the intermediate ester and ring closure, provides the dihydropyrone of formula U-4. Reaction of the dihydropyrone with the aldehyde of formula B-2, prepared as described in Chart B above, in the presence of AlCl<sub>3</sub> provides the benzylidene compound of formula U-5; subsequent reaction with Grignard reagents or trialkyl aluminums in the presence of cuprous bromidedimethyl sulfide complex affords compounds of formula U-6 (wherein, e.g., R<sub>1</sub> is ethyl, tert-butyl, or cyclopropyl). Removal of the benzyloxy-carbonyl (CBZ) protecting group 50 is accomplished using ammonium formate and palladium on charcoal to give the amines of formula U-7 (wherein, e.g., R<sub>1</sub> is ethyl, tert-butyl, or cyclopropyl). Treatment of the amines with sulfonyl chlorides and pyridine in methylene chloride provides the sulfonamides of formula U-8 (wherein, e.g., R<sub>1</sub> is ethyl, tert-butyl, or cyclopropyl and R<sub>2</sub> is alkyl, aryl, or heteroaryl).

#### CHART V

Commercially available 4-fluorobenzaldehyde of formula 60 V-1 is condensed with acetone, under basic conditions, to provide 1,5-Bis-(4-fluorophenyl)-penta-1,4-dien-3-one of formula V-2. The dienone is reduced with magnesium in methanol to provide the ketone of formula V-3. The ketone of formula V-3 is converted to dihydropyrone products of formula V-8 using chemistry analogous to that described in Chart U for the sequence of reactions from U-3 to U-8.

Commercially available trans 2-pentenoic acid of formula W-1 is converted to the corresponding acid chloride using oxalyl chloride in methylene chloride to afford the product of formula W-2. The lithium amide of formula W-3, readily available from the treatment of commercially available (S)-(+)-4-phenyl-2-oxazolidinone with n-butyl lithium in tetrahydrofuran at -78° C., is treated with the acid chloride of formula W-2, to give the unsaturated amide of formula W-4. Addition of the amide of formula W-4 to a tetrahydrofuran solution containing commercially available CuBr/ (CH<sub>3</sub>)<sub>2</sub>S and 3-[bis(trimethylsilyl)amino]phenylmagnesium chloride at -20° C. affords the compound of formula W-5 upon acid workup (Hruby et al., J. Org. Chem., 58(26):7567 (1993)). Treatment of the aniline of formula W-5 with benzyl bromide and sodium carbonate in a water/methylene chloride mixture at reflux; or, potassium carbonate in refluxing acetonitrile, affords the compound of formula W-6. Treatment of the amide of formula W-6 with TiCl<sub>4</sub> followed by an amine base in a solvent such as methylene chloride at below -20° C., preferably at -78° C., then addition of the 2-methoxy-2-methyl-1,3-dioxoline of formula W-7 (prepared as described in Santry et al., J. Am. Chem. Soc., 110(9):2910 (1988)) affords the compound of formula W-8. Brief treatment of the compound of formula W-8 with a protic acid affords the β-ketoamide of formula W-9. Further treatment of the compound of formula W-9 with TiCl<sub>4</sub> followed by an amine base, then 4-heptanone or propylphenethylketone, affords the compound of formula W-10 wherein  $R_1$  is n-propyl or phenethyl, respectively. Treatment of the compound of formula W-10 with sodium hydride or preferably potassium t-butoxide, in an ether solvent then affords the pyrone of formula W-11. Hydrogenation of the compound of formula W-11 using, e.g., a Pd on carbon as the catalyst, affords the compound of formula W-12. Finally, treatment of the compound of formula W-12 with a sulfonyl chloride of formula D-7, wherein R<sub>4</sub> is 5-trifluoromethyl-2-pyridinyl, in an organic solvent, such as methylene chloride, in the presence of an organic base, such as pyridine, provides the final compound of formula W-13, wherein R<sub>1</sub> is n-propyl or phenethyl (when R<sub>1</sub> is phenethyl, it is a pair of diastereomers).

#### CHART X

The final (R) enantiomer of formula X-13, wherein  $R_1$  is n-propyl or phenethyl, is prepared according to the procedures of Chart W.

#### **CHART Y**

Acetyl chloride of formula Y-1 is added to the lithium amide of formula Y-2 (also X-3), readily available from the treatment of commercially available (R)-(-)-4-phenyl-2oxazolidinone with n-butyl lithium in tetrahydrofuran at -78° C., to afford the product of formula Y-3. The compound of formula Y-3 is treated first with TiCl<sub>4</sub> in methylene chloride below room temperature, followed by the addition of a tertiary amine base with subsequent addition of the aldehyde of formula Y-4 (aldehyde of the formula Y-4 is readily available from the reaction of commercially available 3-aminobenzaldehyde with benzyl bromide and potassium or sodium carbonate in either acetonitrile or a water/ methylene chloride mixture) to yield the compound of formula Y-5. Addition of the amide of formula Y-5 to a tetrahydrofuran solution containing commercially available CuBr/(CH<sub>3</sub>)<sub>2</sub>S and ethylmagnesium chloride at -20° C. affords the compound of formula Y-6. Alternatively, the

commercially available compound of formula Y-7 is treated with oxalyl chloride to afford the compound of formula Y-8. The compound of formula Y-8 is then added to a THF solution of the compound of formula Y-2 (also X-3), readily available from the treatment of commercially available (R)-(-)-4-phenyl-2-oxazolidinone with n-butyl lithium in tetrahydrofuran at -78° C., to yield the compound of formula Y-9. Reduction of the compound of formula Y-9 with iron metal in an alcohol/water mixture then affords the compound of formula Y-10. Treatment of the compound of 10 formula Y-10 with benzyl bromide and potassium or sodium carbonate in either acetonitrile or methylene chloride/water then affords the compound of formula Y-5 which, as described above, is converted to the compound of formula Y-6. The compound of formula Y-6 is converted to final 15 product as described for the conversion of the compound of the formula W-6 to the compound of the formula W-13 (wherein R<sub>1</sub> is propyl or phenethyl) in Chart W.

#### CHART Z

Preparation of the (3S) amide of formula Z-6 is accomplished in the same manner as outlined in Chart Y above, except using the compound of formula Z-2 (also W-3). The compound of the formula Z-6 is converted to final product as described for the conversion of the compound of formula 25 X-6 to the compound of the formula X-13 (wherein  $R_1$  is propyl or phenethyl) in Chart Z.

#### CHART AA

Preparation of the 3(S), 6(S) Diastereomers AA-12 and 30 AA-14: Addition of the unsaturated amide of formula AA-1 (also Y-5) to a tetrahydrofuran solution containing commercially available CuBr/(CH<sub>3</sub>)<sub>2</sub>S and ethylmagnesium chloride at -20° C. affords the compound of formula AA-2 (same as Y-6). Reduction of the compound of formula AA-2 with a 35 metal hydride (sodium borohydride, lithium aluminum hydride) affords the compound of formula AA-3. Oxidation of the compound of formula AA-3 (Swern oxidation) affords the aldehyde of formula AA-4 which is treated with trimethylsilylcyanide to yield the trimethylsilyl protected cyano- 40 hydrin of formula AA-5. Alternatively, the compound of formula AA-2 is treated with trimethyl aluminum followed by N-methyl-O-methyl hydroxyl amine to yield the amide of formula AA-6 which is treated with lithium aluminum hydride to yield the aldehyde of formula AA-4. The trim- 45 ethylsilyl cyanohydrin of formula AA-5 is reacted with a strong base (e.g. n-butyl lithium) followed by the addition of chiral epoxide of formula AA-7 (also BB-12; the synthesis of which is described in Chart BB) to yield the compound of formula AA-8. The compound of formula AA-8 is dissolved 50 in methylene chloride and cooled to -78° C. and TiCl<sub>4</sub> is added followed by a tertiary amine base. To that solution is added trimethylorthoformate followed by additional TiCla which yields the compound of formula AA-9. Treatment of the compound of formula AA-9 with base followed by 55 trimethylsilyl chloride, then treatment with an oxidizing agent (ozone), followed by treatment with tetrabutyl ammonium fluoride and then either potassium tert. butoxide or sodium hydride in an ether solvent, then affords the compound of formula AA-10. Hydrogenation of the compound 60 of formula AA-10 then affords the compound of formula AA-11. Finally, treatment of the compound of formula AA-11 with a sulfonyl chloride of formula D-7 in Chart D, wherein R<sub>4</sub> is, e.g., 5-trifluoromethyl-2-pyridinyl, in an organic solvent, such as methylene chloride, in the presence 65 of an organic base, such as pyridine, provides the final compound of formula AA-12.

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Furthermore, addition of the compound of formula AA-1 to a tetrahydrofuran solution containing commercially available CuBr/(CH<sub>3</sub>)<sub>2</sub>S and tertiary butylmagnesium chloride at -20° C. affords the compound of formula AA-13. The compound of formula AA-13 is converted to the final product, the compound of formula AA-14, using the chemistry described for the synthesis of AA-12.

#### CHART BB

Chart BB describes the asymmetric synthesis of epoxides of formula BB-7 and BB-12. Alkylation of 2-methyl-2propen-1-ol (BB-1) with commercially available benzyl bromide provides the allylic alcohol of formula BB-2 (see Lipshutz, B. H. et al.; Synthesis 1992, 191). Catalytic Sharpless epoxidation using commercially available (+) diethyl L-tartrate provides the epoxy alcohol of formula BB-8 (see: (a) Pfenniger, A.; Synthesis 1986, 89. (b) Johnson, R. A.; Sharpless, K. B. In Catalytic Asymmetric Synthesis; Ojima, I., Ed.; VCH: New York, 1993; Chapter 4.1, 103.). Alkylation of the compound of formula BB-8 with benzyl bromide (see: Lipshutz, B. H. et al.; Synthesis 1992, 191) gives the compound of formula BB-9. Reaction of the compound of formula BB-9 with commercially available ethylmagnesium bromide affords the tertiary alcohol of formula BB-10 (see: Hanson, R. M. Chem. Rev. 1991, 91, 437). Catalytic hydrogenolysis of the compound of formula BB-10 provides the diol of formula BB-11. The compound of formula BB-11 is converted to the chiral epoxide of formula BB-12 by standard methodology (for a discussion of the conversion of vicinal diols to epoxides see: Mitsunobu, O. In Comprehensive Organic Synthesis; Trost, B. M. Ed.; Pergamon Press: Oxford, 1991; Vol. 6; Chapter 1.1, 1).

In an analogous manner, the epoxide of formula BB-7 is ultimately derived from the epoxy alcohol of formula BB-3, which in turn is prepared by Sharpless epoxidation of allylic alcohol BB-2 using commercially available (-) diethyl D-tartrate.

Alternatively, reaction of the epoxy alcohol of formula BB-8 with commercially available 4-toluenesulfonyl chloride under standard conditions affords the tosylate of formula BB-13. Reaction of the compound of the formula BB-13 with ethylmagnesium bromide under conditions similar to those described for the nucleophilic opening of arenesulfonate derivatives of glycidol (see: Klunder, J. M.; Onami, T.; Sharpless, K. B. J. Org. Chem. 1989, 54, 1295) affords a mixture of the desired epoxide of formula BB-12 and hydroxytosylate of formula BB-14. The hydroxytosylate of formula BB-14 is readily converted to epoxide BB-12 by the action of  $K_2CO_3$  in methanol.

## CHART CC

Preparation of the 3(S), 6(R) Diastereomers CC-12 and CC-14: These diastereomers are prepared in a manner identical to that described in Chart AA with the exception that the epoxide of formula CC-7 (same as BB-7) is used.

#### CHART DD

Preparation of the 3(R), 6(S) Diastereomers DD-12 and DD14: These diastereomers are prepared in a manner identical to that described in Chart AA with the exception that the amide of formula DD-1 (same as Z-5) is used.

## CHART EE

Preparation of the 3(R), 6(R) Diastereomers EE-12 and EE-14: These diastereomers are prepared in a manner iden-

tical to that described in Chart DD with the exception that the epoxide of formula EE-7 (same as BB-7) is used.

#### CHART FF

The lithium amide of formula FF-2, readily available from the treatment of commercially available (S)-(+)-4phenyl-2-oxazolidinone with n-butyl lithium in tetrahydrofuran at -78° C., is treated with acetyl chloride of formula FF-1 to give the amide of formula FF-3. Treatment of the compound of formula FF-3 with TiCl4 followed by treatment with a trialkyamine followed by the addition of commercially available trimethylacetaldehyde affords the compound of formula FF-4. Addition of the amide of formula FF-4 to a tetrahydrofuran solution containing commercially available CuBr/(CH<sub>3</sub>)<sub>2</sub>S and 3-[bis(trimethylsilyl)amino] phenylmagnesium chloride at -20° C. affords the compound of formula FF-5 upon acid workup. Treatment of the aniline of formula FF-5 with benzyl bromide and sodium carbonate in a water/methylene chloride mixture at reflux; or, potassium carbonate in refluxing acetonitrile, affords the compound of formula FF-6.

The lithium amide of formula FF-7, readily available from the treatment of commercially available (S)-(-)-4benzyl-2-oxazolidinone with n-butyl lithium in tetrahydrofuran at -78 C., is treated with acetyl chloride of formula FF-1 to give the amide of formula FF-8. Treatment of the compound of formula FF-8 with TiCl<sub>4</sub> followed by treatment with a trialkyamine followed by the addition of commercially available trimethylacetaldehyde affords the compound of formula FF-9. Addition of the amide of formula FF-9 to a tetrahydrofuran solution containing commercially available CuBr/(CH<sub>3</sub>)<sub>2</sub>S and 3-[bis(trimethylsilyl)amino] phenylmagnesium chloride at -20° C. affords a mixture of compounds of formulae FF-10a and FF-10b. Treatment of 35 the aniline of formula FF-10b with benzyl bromide and sodium carbonate in a water/methylene chloride mixture at reflux; or, potassium carbonate in refluxing acetonitrile, affords the compound of formula FF-11 Treatment of the compound of formula FF-11 with TiCl<sub>4</sub> in methylene chlo-40 ride followed by the addition of a tertiary amine base then addition of 2-methyl-2-methoxy-1,3-dioxolane affords an intermediate dioxolane (see W-8 in Chart W) which is treated with mild acid to give the compound of formula FF-12. Treatment of the compound of formula FF-12 with TiCl<sub>4</sub>, then a tertiary amine base, followed by addition of either 4-heptanone or 1-phenyl-3-hexanone, affords the aldol product of formula FF-13. Treatment of the compound of formula FF-13 with either sodium hydride or potassium tert. butoxide in an ether solvent then affords the compound of formula FF-14. The compound of formula FF-14 is then hydrogenated under an atmosphere of hydrogen in the presence of a Pd on carbon catalyst to give the compound of formula FF-15. Finally, treatment of the compound of formula FF-15 with a sulfonyl chloride of formula D-7 in Chart D, wherein R<sub>4</sub> is, e.g., 5-trifluoromethyl-2-pyridinyl, in an organic solvent, such as methylene chloride, in the presence of an organic base, such as pyridine, provides the final compound of formula FF-16, wherein R<sub>1</sub> is, e.g., propyl or phenethyl.

#### **CHART GG**

Intermediate of formula GG-6 and final products of formula GG-16 are prepared as described in Chart FF with the exception that the (R)-(-)-4-phenyl-2-oxazolidinone and 65 the (R)-(+)-4-benzyl-2-oxazolidinone chiral auxiliaries are used.

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## CHART HH

The compound of formula HH-1 (W-6), prepared as described in Chart W, is converted to the ester of formula HH-2 wherein R is t-Bu by addition of potassium t-butoxide to a solution of the compound of formula HH-1 in tetrahydrofuran at 0° C. The compound of formula HH-2 wherein R is t-Bu may also be prepared from HH-1 in two steps. First, the oxazolidinone group is cleaved by treatment of the compound of formula HH-1 with lithium hydroxide and hydrogen peroxide at 0° C. in tetrahydrofuran and water. Next, the acid intermediate is treated with N,Ndimethylformamide t-butylacetal in refluxing benzene to produce the ester of formula HH-2 (R is t-Bu). The ester of formula HH-2 wherein R is Me is prepared by heating a mixture of titanium tetrachloride and HH-1 in methanol. The compound of formula HH-3 is prepared by treatment of the ester of formula HH-2 with lithium diisopropylamide or sodium hexamethyldisilylazide to form an enolate, which is then trapped by ethyl formate to give the compound of formula HH-3. Treatment of this intermediate with tosyl chloride in 1,2-dimethoxyethane gives the compound of formula HH-4, which is then converted to the sulfur derivative of formula HH-5 by treatment with a mixture of potassium hydride and thiophenol in tetrahydrofuran. The compound of formula HH-5 is then deprotonated using t-butyllithium in tetrahydrofuran at low temperature. Addition of the epoxide of formula HH-6 (BB-7), prepared as described in Chart BB, and an equivalent of boron trifluoride diethyl etherate affords the compound of formula HH-7. This intermediate is cyclized to the compound of formula HH-8 in situ, or it is isolated and treated with sodium hydride in tetrahydrofuran to produce the cyclic compound of the formula HH-8. The sulfur group is then hydrolyzed using either sodium hydroxide in acetonitrile or aqueous copper chloride to give the dihydropyrone derivative of formula HH-9. The benzyl protecting groups are then removed by catalytic hydrogenation using 10% palladium on carbon in ethyl acetate. The resulting amine of formula HH-10 is converted to the desired sulfonamide derivative of formula HH-11 by treatment with 5-cyanopyridine-2sulfonyl chloride, prepared using the methods described in Chart O, and pyridine in dichloromethane.

#### CHARTS II-00

The diastereomer of formula II-7 is prepared according to Chart II by procedures analogous to those described for the preparation of the diastereomeric product in Chart HH. Likewise, stereoisomers of formulae JJ-11, KK-7, LL-11, MM-7, NN-11, and OO-7 are prepared according to Charts JJ, KK, LL, MM, NN, and OO, respectively, by procedures analogous to those described in Chart HH.

#### CHART PP

The compound of formula PP-4 (HH-8) is also generated as described in Chart PP. The acid of formula PP-2 is prepared by treatment of the t-butyl ester of formula PP-1 (HH-5), prepared as described in Chart HH, with aqueous acid. The compound of formula PP-2 is then treated with t-butyllithium in tetrahydrofuran at low temperature to produce a dianionic intermediate, which is treated with the epoxide of formula PP-3 (BB-7), prepared as described in Chart BB, and an equivalent of boron trifluoride diethyl etherate to afford the compound of formula PP-4 (HH-8).

#### CHARTS QQ-WW

The diastereomer of formula QQ-3 (II-4) is prepared according to Chart QQ by procedures analogous to those

described for the preparation of the diastereomeric product in Chart PP. Likewise, stereoisomers of formulae RR-4 (JJ-8), SS-3 (KK-4), TT-4 (LL-8), UU-3 (MM-4), VV-4 (NN-8), and WW-3 (OO-4) are prepared according to Charts RR, SS, TT, UU, VV, and WW, respectively, by procedures 5 analogous to those described in Chart PP.

#### CHART XX

The compound of formula XX-6 (HH-9) is also generated as described in Chart XX. The compound of formula XX-1 (HH-2), prepared as described in Chart HH, is heated neat in commercially-available tris(dimethylamino)methane, bis (dimethylamino)-methoxymethane or t-butoxy-bis (dimethylamino)methane to generate the intermediate of formula XX-2. One equivalent of t-butyllithium is added to a solution of this ester in tetrahydrofuran at low temperature to produce an anionic intermediate, which is treated with the epoxide of formula XX-3 (BB-7), prepared as described in Chart BB, and an equivalent of boron trifluoride diethyl etherate to afford the compound of formula XX-4. The intermediate of formula XX-4 is cyclized to the dihydropyrone intermediate XX-5 in situ, or XX-4 is isolated and cyclized by treatment with potassium t-butoxide or sodium hydride in tetrahydrofuran. Likewise, intermediate XX-5 is hydrolyzed in situ to form the compound of formula XX-6 (HH-9), or it is isolated and converted to the dihydropyrone of formula XX-6 (HH-9) by treatment with aqueous acid or aqueous base.

#### **CHARTS YY-EEE**

The diastereomer of formula YY-5 (II-5) is prepared according to Chart YY by procedures analogous to those described for the preparation of the diastereomeric product in Chart XX. Likewise, stereoisomers of formulae ZZ-6 35 (JJ-9), AAA-5 (KK-5), BBB-6 (LL-9), CCC-5 (MM-5), DDD-6 (NN-9), and EEE-5 (OO-5) are prepared according to Charts ZZ, AAA, BBB, CCC, DDD, and EEE, respectively, by procedures analogous to those described in Chart XX.

#### **CHART FFF**

The diastereomers of formulae FFF-5 and FFF-7 are also prepared by separation of a diastereomeric intermediate. The diastereomeric mixture of formula FFF-1 (W-11), prepared as described in Chart W, is separated into the single diastereomers of formulae FFF-2 (less polar diastereomer) and FFF-3 (more polar diastereomer) using a preparative chiral HPLC column. The benzyl protecting groups of compounds FFF-2 and FFF-3 are then removed by catalytic hydrogenation using 10% palladium on carbon in ethyl acetate to form the amines of formulae FFF-4 and FFF-6, respectively. The amine intermediates are then converted to the desired sulfonamide derivatives of formulae FFF-5 (HH-11) and FFF-7 (II-7), respectively, by treatment with 5-cyanopyridine-2-sulfonyl chloride, prepared using the methods described in Chart O, and pyridine in dichloromethane.

#### **CHART GGG**

The m-nitrocinnamic acid chloride (available from the treatment of the commercially available acid with oxylal chloride) of formula GGG-1 is added to an ether solution of the lithiooxazolidinone of formula GGG-2 (readily available 65 from the treatment of commercially available (R)-(+)-4-benzyl-2-oxazolidinone with n-butyl lithium) to afford the

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compound of formula GGG-3. The compound of formula GGG-3 is treated with either SnCl<sub>2</sub>.2H<sub>2</sub>O in ethanol or iron powder in a mixture of ethanol/water and containing ammonium chloride, to effect the reduction of the nitro group to the corresponding amine found in the compound of formula GGG-4. The compound of formula GGG-4 is treated with excess benzyl bromide in the presence of potassium or sodium carbonate in an organic solvent (with methylene chloride/water also being added) to yield the compound of formula GGG-5. Addition of a THF solution of the compound of formula GGG-5 to a THF/dimethylsulfide mixture containing the cuprate reagent prepared from ethyl magnesium bromide and copper bromide/dimethyl sulfide complex affords the compound of formula GGG-6. The compound of GGG-6 is then treated with TiCl<sub>4</sub>, then a tertiary amine, followed by the addition of 2-methyl-2-methyoxy-1,3dioxolane of formula GGG-7 to yield the compound of formula GGG-8. Treatment of the compound of formula GGG-8 with perchloric acid then yields the compound of formula GGG-9. Alternately, the compound of formula GGG-6 is treated with a strong base such lithium diisopropylamide in an ether solvent below room temperature and added to a solution of acetyl chloride (also in an ether solvent and cooled to below room temperature) to yield the compound of formula GGG-9. The compound of formula GGG-9 is treated with TiCl<sub>4</sub> in methylene chloride followed by the addition of a tertiary amine, then addition of either 4-heptanone or 1-phenyl-3-hexanone to yield the compound of formula GGG-10. The compound of formula GGG-10 is then treated with either sodium hydride or potassium tertbutoxide in an ether solvent to yield the compound of formula GGG-11. The compound of formula GGG-11 is then hydrogenated to yield the compound of formula GGG-12. The compound of formula GGG-12 is then converted to the final title compound by treatment with a sulfonyl chloride of formula D-7 in Chart D, wherein R<sub>4</sub> is, e.g., 5-trifluoromethyl-2-pyridinyl, in an organic solvent, such as methylene chloride, in the presence of an organic base, such as pyridine, provide the final compound of formula GGG-13, wherein  $R_1$  is, e.g., n-propyl or phenethyl.

Alternatively, addition of the compound of formula GGG-5 to a THF/dimethylsulfide solution containing a mixture of tert-butyl magnesium chloride and copper bromide/dimethylsulife complex at below 0° C. yields a mixture of compounds of formulae GGG-14a and GGG-14b. Both the compounds of formulae GGG-14a and GGG-14b are converted to the final products GGG-19 and GGG-20 using the methodology described in Chart GGG for the synthesis of the C-3 ethyl compound of formula GGG-13.

#### **CHART HHH**

The final compounds of formula HHH-13, HHH-19 and HHH-20 are prepared in the same manner as described for the final compounds in Chart GGG.

## CHART III

The commercially available acid of formula III-1 is converted to the compound of formula III-2 by treatment with oxalyl chloride. The acid chloride of formula III-3 is then coupled to the lithio oxazolidinone of formula III-3 (readily available from the treatment of commercially available (S)-(-)-4-benzyl-2-oxazolidinone with n-butyl lithium in an ether solvent) to yield the compound of formula III-4. Addition of the amide of formula III-4 to a tetrahydrofuran solution containing commercially available copper bromide/dimethyl sulfide complex and 3-[bis(trimethylsilyl)amino]

phenylmagnesium chloride at -20° C. affords the compounds of formula III-5a and III-5b upon acid workup. These compounds are separable by silica gel chromatography. The compound of formula III-5a is treated with benzyl bromide in either acetonitrile or a methylene chloride/water 5 mixture in the presence of either potassium or sodium carbonate to yield the compound of formula III-6. The compound of formula III-6 is treated with TiCl4 in methylene chloride followed by the addition of a tertiary amine and then 2-methyl-2-methoxy-1,3-dioxolane of formula III-7 is 10 added to yield the compound of formula III-8. Treatment of the compound of the formula III-8 with an acid such as perchloric acid then yields the compound of formula III-9. Treatment of the compound of formula III-9 with TiCla in methylene chloride then addition of a tertiary amine, fol- 15 lowed by the addition of either 4-heptanone or 1-phenyl-3hexanone then affords the compound of formula III-10. Treatment of the compound of formula III-10 with either sodium hydride or potassium tert. butoxide then affords the compound of formula III-11. The compound of formula 20 III-11 is hydrogenated to afford the compound of formula III-12. Finally, treatment of the compound of formula III-12 with a sulfonyl chloride of formula D-7 in Chart D, wherein R<sub>4</sub> is, e.g., 5-trifluoromethyl-2-pyridinyl, in an organic solvent, such as methylene chloride, in the presence of an 25 organic base, such as pyridine, provides the final compound of formula III-13, where in R<sub>1</sub> is, e.g., propyl or phenethyl.

In an analogous fashion, starting with the compound of formula III-5b, the final compound of formula III-14 is also prepared.

#### CHART JJJ

The final compounds of formula JJJ-13 and JJJ-14 are prepared using the methodology described in Chart III.

#### CHART KKK

The compound of formula KKK-1 (same as JJJ-9) is treated with TiCl<sub>4</sub> in methylene chloride followed by the addition of a tertiary amine. To that solution is added commercially available hydrocinnamaldehyde to afford the compound of formula KKK-2. The compound of formula KKK-2 is oxidized (e.g. Me<sub>2</sub>SO—SO<sub>3</sub>/pyridine) to yield the compound of formula KKK-3. The compound of formula KKK-3 is treated with propylmagnesium chloride (where R<sub>1</sub> 45 is, e.g., phenyl) to yield the compounds of formula KKK-4a and KKK-4b. Depending on the specific reaction conditions, the ratio of KKK-4a/KKK-4b varies. Alternatively, addition of allylzinc bromide or allylsilane in the presence of TiCl, or n-Bu<sub>4</sub>NF (see Taniguchi et. al. Chemistry Letters 2135, 50 1992) to the compound of formula KKK-3, followed by hydrogenation, also yields the compounds of formula KKK-4a and KKK-4b. Depending on the specific reaction conditions the ratio of KKK-4a and KKK-4b vary. The compound of KKK-4a is treated with either sodium hydride or potas- 55 sium tert, butoxide to yield the compound of formula KKK-5. It is also possible that upon treatment of KKK-3 with allyl zinc bromide, allyl silane or propylmagnesium chloride the intermediate metal alkoxide (metals being magnesium, zinc and titanium) will undergo spontaneous 60 cyclization to yield an unsaturated intermediate which upon hydrogenation leads directly to KKK-5 without the isolation of KKK-4a. The compound of formula KKK-5 is hydrogenated to yield the compound of formula KKK-6. Finally, treatment of the compound of formula KKK-6 with a 65 sulfonyl chloride of formula D-7 in Chart D, wherein R4 is, e.g., 5-trifluoromethyl-2-pyridinyl, in an organic solvent,

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such as methylene chloride, in the presence of an organic base, such as pyridine, provides the final compound of formula KKK-7a, wherein, e.g.,  $R_1$  and  $R_2$  are phenyl or propyl, respectively.

In an analogous manner to that described for the conversion of the compound of formula KKK-4a to the compound of formula KKK-7a, the compound of formula KKK-4b is converted to the final product of formula KKK-7b.

In an analogous manner to that described for the conversion of the compound of formula KKK-1 to final products of the formula KKK-7a and KKK-7b, the compounds of formula KKK-14a and KKK-14b, wherein  $R_1$  and  $R_2$  are, e.g., methyl or phenethyl, respectively, are prepared by starting with the compound of formula KKK-8 (same as III-6).

In an analogous manner to that described for the conversion of the compound of formula KKK-1 and the compound of formula KKK-8 (each containing the 4-benzyl-2-oxazolidinone auxillary) to the final products of the formulae KKK-7a and KKK-7b, and the final formulae KKK-14a and KKK-14b respectively, the compounds of the formula KKK-15 and the compound of the formula KKK-19 (each containing the 4-phenyl-2-oxazolidinone auxillary) are converted to the final products of the formula KKK-7a and K-7b, and the final products of formula KKK-14a and KKK-14b, respectively, wherein R<sub>1</sub> and R<sub>2</sub> are, e.g., methyl or phenethyl, respectively.

#### CHART LLL

The compound of formula LLL-1 (same as: wherein R is phenyl, AA-1; wherein R is benzyl, GGG-5) is added to a THF solution of commercially available copper bromide/ dimethylsulfide complex and tert. butylmagnesium chloride below 0° C. to afford the compound of formula LLL-2 as the major diasteromeric product. Where R is defined as benzyl 35 in the compound of formula LLL-2, that compound is treated with TiCl<sub>4</sub> in methylene chloride below 0° C. followed by the addition of a tertiary amine, then the addition of 2-methyl-2-methoxy-1,3-dioxolane to yield the compound of formula LLL-3. The compound of formula LLL-3 is treated with a protic acid to afford the compound of formula LLL-4. The compound of formula LLL-4 is treated with TiCl<sub>4</sub> in methylene chloride below 0° C. followed by the addition of an amine base, then addition of either 4-heptanone or 1-phenyl-3-hexanone affords the compound of formula LLL-5 wherein R, is, e.g., n-propyl or phenethyl, respectively. Treatment of the compound of formula LLL-5 with either sodium hydride or potassium tert, butoxide in an ether solvent affords the pyrone of formula LLL-6. Hydrogenation of the compound of formula LLL-6 using, e.g. a Pd on carbon as the catalyst, affords the compound of formula LLL-7. Finally, treatment of the compound of formula LLL-7 with a sulfonyl chloride of formula D-7 in Chart D, wherein R<sub>4</sub> is, e.g., 5-trifluoromethyl-2-pyridinyl, in an organic solvent, such as methylene chloride, in the presence of an organic base, such as pyridine, provides the final compound of formula LLL-8, wherein R<sub>1</sub> is, e.g., propyl or phenethyl.

The compound of formula LLL-2, where R is phenyl, is treated with TiCl<sub>4</sub> in methanol to yield the compound of formula LLL-9. The compound of formula LLL-9 is treated with a base to effect hydrolysis to give the compound of formula LLL-10. The acid of formula LLL-10 is treated with methyl lithium in an ether solvent to yield the compound of formula LLL-11. The ketone of formula LLL-11 is treated with TiCl<sub>4</sub> in methylene chloride below 0° C. followed by the addition of an amine base, then addition of either

4-heptanone or 1-phenyl-3-hexanone, to give the compound of formula LLL-12 wherein R<sub>1</sub> is, e.g., n-propyl or phenethyl, respectively. The compound of formula LLL12 is treated with TiCl<sub>4</sub> in methylene chloride below 0° C. followed by the addition of an amine base, then the addition of trimethyl orthoformate to yield the compound of formula LLL-13. The compound of formula LLL-13, in an organic solvent such as THF or methylene chloride, is treated with a base followed by the addition of trimethylsilyl chloride. The solvent is removed from the aforementioned reaction 10 and the resulting protected tertiary alcohol is oxidized (e.g. Ru cat./t-BuOH (see Murahashi et. al. Chemistry Letters 2237, 1992); tritylperchlorate/methylene chloride (see Mukaiyama et. al. Chemistry Letters 1255, 1985), ozone/ methylene chloride (see Can. J. Chem. 49, 2465, 1971)) to 15 dichloromethane. afford the lactone LLL-6 directly or in a two step sequence where the intermediate ester is lactonized with the aid of either sodium hydride, potassium tert. butoxide or n-Bu<sub>4</sub>NF in an ether solvent. The conversion of the compound of formula LLL-6 to the final product is described above.

Following the same strategy the compound of formula LLL-16 is converted to the final products of formula LLL-23 wherein  $R_1$  is propyl or phenethyl.

#### **CHART MMM**

The diastereomers of formulae MMM-5 and MMM-7 are also prepared by separation of a diastereomeric mixture of these two compounds. Alternatively, the diastereomeric mixture of formula MMM-1 (X-11 where  $R_1$  is, e.g.,  $_{30}$ phenethyl) prepared as described in Chart X, is separated into the single diastereomers of formulae MMM-2 and MMM-3 using a preparative chiral HPLC column. The benzyl protecting group groups of compounds MMM-2 (less polar diastereomer) and MMM-3 (more polar diastereomer) 35 are then removed by catalytic hydrogenation using 10% palladium on carbon in ethyl acetate to form the amines of formulae MMM-4 and MMM-6, respectively. The amine intermediates are then converted to the desired sulfonamide derivatives of formulae MMM-5 and MMM-7, respectively, 40 by treatment with 5-trifluoromethyl-2-pyridinylsulfonyl chloride, prepared using the methods described in Chart O, and pryidine in methylene chloride.

#### **CHART NNN**

The commercially available (1R,2S)-(-)ephedrine of formula NNN-2 is treated with triethylamine and the acid chloride of formula NNN-1 (W-2), prepared as described in Chart W, to afford the amide of formula NNN-3. A t-butvl methyl ether solution of this amide at 0° C. is treated 50 sequentially with 1.1 equivalents of propyl magnesium chloride and 2.0 equivalents of 3-[bis(trimethylsilyl)amino] phenyl magnesium chloride, stirred for 3 hours at 0° C., washed with ammonium chloride solution and concentrated in vacuo. The residue is then stirred with silica gel in 55 chloroform to afford the compound of formula NNN-4. Alternatively, the above reaction mixture may be washed with 1N hydrochloric acid solution during the workup instead of ammonium chloride solution to generate the compound of formula NNN-4. The amine is then converted 60 to the derivative of formula NNN-5 by heating a mixture of the compound of formula NNN-4, 2.2 equivalents of benzyl bromide and 2.2 equivalents of sodium carbonate in acetonitrile. The intermediate of formula NNN-5 is then treated with 2 equivalents of lithium diisopropylamide in tetrahy- 65 drofuran to form the lithium enolate, which is trapped with acetyl chloride to afford the b-ketoamide of formula NNN-6.

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A solution of this amide in methylene chloride at low temperature may be treated with 1 equivalent of titanium tetrachloride and 1 equivalent of diisopropylethylamine, followed by 4-heptanone to generate the compound of formula NNN-7. Conversion of the amide of formula NNN-7 to the dihydropyrone of formula NNN-8 may be accomplished with either sodium hydride in tetrahydrofuran or with aqueous acid. The benzyl protecting groups may then be removed by catalytic hydrogenation using 10% palladium on carbon in ethyl acetate. The resulting amine of formula NNN-9 is converted to the desired sulfonamide derivative of formula NNN-10 (W-12) by treatment with 5-trifluoromethylpyridine-2-sulfonyl chloride, prepared using the methods described in Chart O, and pyridine in dichloromethane.

#### CHART 000

The compound of formula OOO-7 (NNN-8) may also be generated as described in Chart OOO. The amide of formula OOO-1 (NNN-5) is treated with aqueous acid to afford the compound of formula OOO-2. The methyl ester of formula OOO-3 is formed from the compound of formula OOO-2 using catalytic acid in methanol. Treatment of the methyl ester of formula OOO-3 with lithium diisopropylamide, followed by trimethylsilyl chloride gives the compound of formula OOO-4. Treatment of this intermediate with either 2-methoxy-2-methyl-1,3-dioxolane followed by hydrolysis or treatment with acetyl chloride affords the β-keto ester of formula OOO-5. This β-keto ester is converted to the compound of formula OOO-6 by treatment of either the titanium enolate (formed using 1 equivalent of titanium tetrachloride and 1 equivalent of diisopropylethylamine in methylene chloride at low temperature) or the lithium dianion (formed using 2 equivalents of lithium diisopropylamide in tetrahydrofuran at low temperature) with 4-heptanone. The dihydropyrone of formula OOO-7 (NNN-8) is formed by treatment of the compound of formula OOO-6 with either sodium hydride in tetrahydrofuran or aqueous base.

## CHART PPP

Reduction of the commercially available ethyl 4,4,4-trifluorobutyrate of formula PPP-1, with DiBAL-H followed by in situ alkylation with 2-phenethyl magnesium bromide or chloride produces the alcohol of formula PPP-2. Swern oxidation of the alcohol gives the ketone of formula PPP-3. The ketone is converted to the dihydropyrone of formula PPP-4 by alkylation with the diamion of methyl acetoacetate followed by saponification to the acid and lactonization with base.

#### **CHART QQQ**

The aluminum trichloride catalyzed reaction of the dihydropyrone of formula QQQ-1 (PPP-4), prepared as described in Chart PPP, with the CBZ-protected 3-aminobenzaldehyde (which is available from the reaction of benzyl chloroformate with commercially available 3-aminobenzaldehyde) of formula QQQ-2 and subsequent reaction with trialkyl aluminums or Grignard reagents in the presence of cuprous bromide-dimethylsulfide complexes provides compounds of formula QQQ-3. The individual stereoisomers are separated by HPLC using a chiral stationary phase to give the four possible stereoisomers of formula QQQ-4, QQQ-5, QQQ-6, and QQQ-7. Transfer hydrogenation of each stereoisomer with Pd/C and ammonium formate gives the amines of formula QQQ-8, QQQ-9, QQQ-10, and QQQ-11. Treatment of the amines with sulfonyl chlorides of

general formula QQQ-12 and pyridine in methylene chloride provides compounds of general formula QQQ-13, QQQ-14, QQQ-15, and QQQ-16, wherein R<sub>2</sub> is, e.g., 5-cyano-2pyridinyl, 1-methyl-4-imidazolyl, or 5-amino-2-pyridinyl.

#### **CHART RRR**

The procedure for the preparation of compounds of formula RRR-11 to RRR-15 is described in Chart RRR. The pyrone RRR-A is coupled to the Cbz protected benzaldehyde RRR-B in THF with AlCl<sub>3</sub> followed by treatment of the resulting intermediate with R<sub>1</sub>MgX where X=Br or Cl in THF with added CuBr.Me<sub>2</sub>S to give RRR-1. De-protection of the resulting intermediate with 10% Pd/C in methanol with added ammonium formate gives RRR-2. Separation of the racemic compound RRR-1 into its 4 enantiomers gives RRR-3 to RRR-6. De-protection of the resulting intermediates with 10% Pd/C in methanol with added ammonium formate gives the free amines RRR-7 to RRR-10. Treatment of the amines RRR-7 to RRR-10 and RRR-2 with an appropriate sulfonyl chloride gives the sulfonamides RRR-11 to RRR-14 and RRR-15, respectively.

#### **CHART SSS**

mula SSS-7 to SSS-9 is described in Chart SSS. The pyrone SSS-A is coupled to the Cbz protected benzaldehyde SSS-B in THF with AlCl<sub>3</sub> followed by treatment of the resulting intermediate with R<sub>1</sub>MgX where X=Br or Cl in THF with added CuBr.Me<sub>2</sub>S to give SSS-1. De-protection of the 30 resulting intermediate with 10% Pd/C in methanol with added ammonium formate gives SSS-2. Separation of the racemic compound SSS-1 into its 2 enantiomers gives SSS-3 to SSS-4. De-protection of the resulting intermediates with 10% Pd/C in methanol with added ammonium formate gives 35 the free amines SSS-5 to SSS-6. Treatment of the amine with an appropriate sulfonyl chloride gives the sulfonamides SSS-7 to SSS-9.

#### **CHART TTT**

The procedure for the preparation of compounds of formula TTT-6 and TTT-7 is described in Chart TTT. The pyrone TTT-A is coupled to the Cbz protected benzaldehyde TTT-B in THF with AlCl<sub>3</sub> followed by treatment of the resulting intermediate with R<sub>1</sub>MgX where X=Br or Cl in THF with added CuBr.Me<sub>2</sub>S to give TTT-1. Separation of the racemic compound TTT-1 into its 2 enantiomers gives TTT-2 and TTT-3. De-protection of the resulting intermediates with 10% Pd/C in methanol with added ammonium formate gives the free amines TTT-4 and TTT-5. Treatment of the amine with an appropriate sulfonyl chloride gives the sulfonamides TTT-6 and TTT-7.

## **CHART UUU**

Reaction between commercially available thiourea in hot ethanol with commercially available 2-chloro-5nitropyridine of formula UUU-1 affords the isothiourea compound of formula UUU-2. Treatment of the compound of formula UUU-2 with aqueous sodium carbonate and 60 sodium hydroxide provides the thiol compound of formula UUU-3. Oxidation of the compound of formula UUU-3 with chlorine gas provides the sulfonyl chloride compound of formula UUU-4. Treatment of the compound of formula D-5 (e.g., the compound of formula T-4 wherein R<sub>1</sub> is 65 2-phenylethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl) in dichloromethane with two equivalents of pyridine followed by one

equivalent of the compound of formula UUU-4 gives the sulfonamide compound of formula UUU-5 (wherein R<sub>1</sub> is 2-phenylethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl). Reduction of the compound of formula UUU-5 with palladium on carbon and ammonium formate affords the compound of formula UUU-6, which is the compound: 5-amino-N-[3-(1-[5,6dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2Hpyran-3-yl]-2,2-dimethylpropyl)phenyl]-2pyridinesulfonamide (Formula UUU-6: R, is 2-phenylethyl, 10 R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl).

#### **CHART VVV**

The compound of Formula VVV-1, which is 2-mercapto-5-carbamoylpyridine, is prepared via published procedure (J. Chem. Soc. 1948, 1939-1945). Treatment of a suspension of this compound in dilute hydrochloric acid with chlorine gas at 0° provides the sulfonyl chloride of Formula VVV-2.

#### **CHART WWW**

Amines of the generic formula WWW-1 are reacted with benzyl chloroformate to provide CBZ derivatives WWW-2. The individual stereoisomers of formula WWW-2 are gen-The procedure for the preparation of compounds of for- 25 erally separated by chiral HPLC methods, and then converted back to the free amines WWW-3 via hydrogenolysis. Sulfonation of the amines in the usual manner known to one of ordinary skill in the art provides the final compounds of formula WWW-4, in stereochemically pure form.

#### **CHART XXX**

Dihydropyrone XXX-1, which is prepared by procedures analogous to those in described in Preparations 17 and 84, is condensed with meta-nitrobenzaldehyde in the presence of aluminum trichloride to provide the benzylidene intermediate XXX-2. Conjugate reduction of the double bond using sodium cyanoborohydride, followed by reduction of the nitro group via catalytic hydrogenation, affords amine of formula XXX-4, which is converted to the sulfonamides XXX-5 by treatment with the appropriate sulfonyl chloride in dichloromethane and pyridine.

#### **CHART YYY**

Dihydropyrones of Formula YYY-1, wherein R<sub>1</sub> and R<sub>2</sub> are propyl or phenethyl, and which are synthesized as described in Preparation 84, are condensed with the aldehyde of Formula B-2 using aluminum trichloride to provide the benzylidene intermediates of formula YYY-2. Conjugate addition of tert-butylmagnesium chloride in the presence of copper (I) bromide-dimethyl sulfide provides compounds of Formula YYY-3. Hydrogenolytic deprotection affords amines of formula YYY-4, which are converted to the sulfonamides of formula YYY-5 using the appropriate sulfonyl chloride in dichloromethane with added pyridine. The procedures used are analogous to those described for Chart

#### CHART ZZZ

Polymeric meta-aminobenzaldehyde is protected by treating with benzyl bromide and potassium carbonate in acetonitrile at reflux to yield the compound of formula ZZZ-2. A vinyl anion is generated from 2-bromovinyltrimethylsilane of formula ZZZ-3 by treatment with t-butyl lithium at -78° C. to -20° C. The vinyl anion so generated is cooled to -78° C. and the diprotected meta-aminobenzaldehyde of formula ZZZ-2 is added to afford the desired allylic alcohol of

formula ZZZ-3. The alcohol is easily converted to the acetate or carbonate of formula ZZZ-5 by standard means (e.g., CH<sub>3</sub>COCl, pyridine, CH<sub>2</sub>Cl<sub>2</sub>, 0° C.). These substrates participate in palladium catalyzed allylic substitutions as delineated in Charts AAAA-CCCC (C. G. Frost; J. Howarth; J. M. J. Williams, Tetrahedron: Asymmetry (1992) 3:1089–1122).

#### **CHART AAAA**

The sodium salt of methyl acetoacetate of formula 10 AAAA-1 generated by treating methyl acetoacetate with sodium hydride at 0° C. in either DMF or THF acts as the nucleophile in a palladium catalyzed allylic substitution. If this reaction is run in the presence of palladium allyl chloride dimer of formula AAAA-3 as the palladium source 15 and a chiral phosphine ligand (P. von Matt; A. Pfaltz, Angew. Chem. Int. Ed. Engl. (1993) 32:566-568), a kinetic resolution of the starting allylic acetate or carbonate results in the synthesis of optically enriched allylated product of formula AAAA-4. If the reaction with nucleophile is slow, the 20 acetate generated from formation of the pi-allyl palladium intermediate isomerizes the two possible diastereomeric pi-allyl complexes so that a stereoselective synthesis of the allylated product occurs (B. M. Trost; P. E. J. Strege, Am. Chem. Soc. (1977) 99:1649). Treatment of the resulting 25 vinyl silane of formula AAAA-4 with para-toluenesulfonic acid in acetonitrile at reflux affords the desilylated olefin of formula AAAA-5. The dihydropyrone product of formula AAAA-7 is formed by generating the dianion of the β-ketoester under standard conditions (J. R. Peterson; T. J. 30 Winger; C. P. Miller, Syn. Comm. (1988) 18(9):949-963), (NaH, n-butyllithium, THF) and quenching with an appropriate symmetrical ketone of formula AAAA-6 (such as 4-heptanone). Hydrolysis of the ester (0.1N NaOH/THF) and acidic work-up provide the dihydropyrone product of 35 formula AAAA-7. Standard hydrogenation conditions reduce the olefin and deprotect the amine. Subsequent treatment of the amino compound with the appropriate sulfonyl chloride of formula AAAA-8 (pyridine, CH<sub>2</sub>Cl<sub>2</sub>) provides the desired sulfonamide protease inhibitor of for- 40 mula AAAA-9.

#### **CHART BBBB**

Alternatively, the palladium catalyzed allylic substitution may be performed with the sodium anion of the requisite 45 dihydropyrone J. R. (Peterson; T. J. Winger; C. P. Miller, Syn. Comm. (1988) 18(9):949-963) of formula BBBB-1 (dihydropyrone, NaH, THF or DMF, 0° C.) as the nucleophilic partner. Once again, if palladium allyl chloride dimer of formula BBBB-3 and a chiral phosphine ligand (P. von 50 Matt; A. Pfaltz, Angew. Chem. Int. Ed. Engl. (1993) 32:566-568) are employed as catalyst, a kinetic resolution results in the synthesis of optically pure allylated dihydropyrone of formula BBBB-4; and a stereoselective synthesis of the allylated product will occur if the reaction with 55 nucleophile is slow relative to isomerization of the two possible diastereomeric pi-allyl complexes by acetate generated from formation of the pi-allyl palladium intermediate. Subsequent, desilylation (p-TsOH, CH<sub>3</sub>CN), olefin reduction and amine deprotection (H<sub>2</sub>/Pd/C), and sulfonylation of 60 the amine (ArSO<sub>2</sub>Cl, pyridine, CH<sub>2</sub>Cl<sub>2</sub>) with a compound of the formula BBBB-5 provides the desired dihydropyrone protease inhibitor of formula BBBB-6.

### CHART CCCC

Treatment of m-bis(benzyl)aminobenzoic acid of formula CCCC-1 with oxalyl chloride to form the acid chloride and

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reaction with bis(trimethylsilyl)acetylene and AlCl<sub>3</sub> in methylene chloride affords the propargylic ketone of formula CCCC-2. Asymetric reduction of the ketone with a chiral borane (H. C. Brown; Beeraraghavan Ramachandran, P. Acc. Chem. Res. (1992) 25:16-24) such as DIP chloride [(+) or (-)-β-chlorodiisopinocampheylborane] and acetylene reduction with REDAL provides the allylic alcohol of formula CCCC-3, primarily as a single enantiomer. Formation of the carbonate of formula CCCC-4 (methyl chloroformate, pyridine, CH<sub>2</sub>Cl<sub>2</sub>, 0° C.) and subjection to palladium catalyzed allylic substitution with the desired dihydropyrone of formula CCCC-5 as nucleophile affords primarily one enantiomer of the allylated dihydropyrone of formula CCCC-6 (retention of configuration) (T. Hayashi; T. Hagihara; M. Konishi; M. J. Kumada, Am. Chem. Soc. (1983) 105:7768-7770). This product is transformed into the desired protease inhibitor of formula CCCC-7 as previously described in Chart BBBB.

### **CHART DDDD**

The known cycloalkylpyranones of formula DDDD-1 are prepared by acylation of the trimethylsilyl enol ether of the corresponding cycloalkyl ketone with malonyl dichloride as described in R. Effenberger, T. Ziegler, K.-H. Schonwalder, T. Kesmarszky, B. Bauer Chem. Ber. 119:3394-3404 (1986). Catalytic hydrogenation of the cycloalkylpyranones of formula DDDD-1 with platinum oxide (PtO2) in acetic acid produces the cycloalkyldihydropyrones of Formula DDDD-2. The intermediate of formula DDDD-3 is then formed by aluminum chloride (AlCl<sub>3</sub>) catalyzed condensation of the compound of formula DDDD-2 with 3-nitrobenzaldehyde, which is commercially available. Subsequent reaction of the intermediate of formula DDDD-3 with trialkyl aluminums in the presence of copper bromidedimethyl sulfide complex (CuBr-Me2S) or zinc reagents generated from zinc metal, alkyl halide, cuprous cyanide (CuCN) and lithium chloride (LiCl) provides compounds of formula DDDD-4 which contain a C-3a branched substituent. Catalytic hydrogenation of compounds of the formula DDDD-4 with Pd/C in ethanol (EtOH) provides the amine derivatives of the formula DDDD-5. Treatment of the compounds of formula DDDD-5 with sulfonyl chlorides of formula DDDD-6 and pyridine in methylene chloride (CH<sub>2</sub>Cl<sub>2</sub>) provides compounds of the formula DDDD-7 (e.g., wherein n is 1, 2, or 3; R<sub>1</sub> is ethyl or t-butyl; R<sub>2</sub> is 4-cyanophenyl or 5-cyano-2-pyridyl).

Procedures by which the compounds of the present invention are prepared are also described in International application, PCT/US93/10645, filed 9 Nov. 1993 (WO 94/11361, published 26 May 1994), and International application, PCT/US94/00938, filed 3 Feb. 1994 (WO 94/18188, published 18 Aug. 1994), both of which are incorporated by reference herein.

As is apparent to those of ordinary skill in the art, the compounds of the present invention can occur in several diastereomeric forms, depending on the configuration around the asymmetric carbon atoms. All such diastereomeric forms are included within the scope of the present invention.

Also, the dihydropyrones of the present invention can be separated into individual stereoisomers or prepared as individual diastereomers. A diastereomeric pair can be prepared wherein C-3 $\alpha$  is a homogeneous center and C-6 is a mixture. All such enantiomeric and diastereomeric forms, and mixtures thereof, are included within the scope of the present invention.

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The compounds of the present invention of formula I can

exist in several tautomeric forms, including the particular

enol forms as depicted by formula I and IA and the keto form

of formula IB. (For formulas I, IA and IB, the dashed line

indicates that a double bond may be present or absent.) All

such tautomeric forms are included within the scope of the

present invention. For compounds of the present invention

which are 4-hydroxy-pyran-2-ones of formula VII, the enol

form predominates. For compounds of the present invention

which are 5,6-dihydro-4-hydroxy-pyran-2-ones of formula

VI, a mixture of the enol and keto forms is commonly

Also, the compounds of the present invention of formula II can exist in several tautomeric forms of the 4-hydroxy-pyrone ring, including the particular enol forms depicted by formulas II and IIA, and the particular keto form depicted by formula IIB, and mixtures thereof. All such tautomeric forms are included within the scope of the present invention.

The compounds of the present invention may be in either free form or in protected form at one or more of the remaining (not previously protected) carboxyl, amino, hydroxy, or other reactive groups. The protecting groups may be any of those known in the art. Examples of nitrogen and oxygen protecting groups are set forth in T. W. Greene, Protecting Groups in Organic Synthesis, Wiley, New York, (1981); J. F. W. McOmie, ed. Protective Groups in Organic Chemistry, Plenum Press (1973); and J. Fuhrhop and G. Benzlin, Organic Synthesis, Verlag Chemie (1983). Included among the nitrogen protective groups are t-butoxycarbonyl (BOC), benzyloxycarbonyl, acetyl, allyl, phthalyl, benzyl, benzoyl, trityl and the like.

The present invention provides for compounds of formula I and II or pharmacologically acceptable salts and/or hydrates thereof. Pharmacologically acceptable salts refers to those salts which would be readily apparent to a manufacturing pharmaceutical chemist to be equivalent to the parent compound in properties such as formulation, stability, patient acceptance and bioavailability. Examples of salts of the compounds of formula I include acidic salts, such as sodium, potassium, lysine, arginine and calcium salts, and basic salts, such as the hydrochloride salt, wherein the R substituents in formula I contain a basic moiety. Examples of salts of the compounds of formula II include the hydrohalide salts, such as the hydrochloride and hydroiodide salts; and the sodium, potassium, calcium, lysine and arginine salts.

Also included as salts of the compounds of formulae I and II of the present invention are the bis-salts, such as the bis-arginine, bis-lysine, bis-sodium, bis-potassium and bis-calcium salts, provided that the compound contains, for example, —NHSO<sub>2</sub>—, —SO<sub>3</sub>H, —CONH—, —OH or COOH. The bis-sodium salt is most preferred.

The compounds of the present invention are useful for treating patients infected with human immunodeficiency virus (HIV) which results in acquired immunodeficiency syndrome (AIDS) and related diseases. For this indication, these compounds may be administered by oral, intranasal, transdermal, subcutaneous and parenteral (including intramuscular and intravenous) routes in doses of 0.1 mg to 100 mg/kg of body weight per day.

Those skilled in the art would know how to formulate the compounds of this invention into appropriate pharmaceutical dosage forms. Examples of the dosage forms include oral formulations, such as tablets or capsules, or parenteral formulations, such as sterile solutions.

When the compounds in this invention are administered orally, an effective amount is from about 0.1 mg to 100 mg

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per kg of body weight per day. Either solid or fluid dosage forms can be prepared for oral administration. Solid compositions, such as compressed tablets, are prepared by mixing the compounds of this invention with conventional ingredients such as talc, magnesium stearate, dicalcium phosphate, magnesium aluminum silicate, calcium sulfate, starch, lactose, acacia, methyl cellulose, or functionally similar pharmaceutical diluents and carriers. Capsules are prepared by mixing the compounds of this invention with an appropriately sized hard gelatin capsule. Soft gelatin capsules are prepared by machine encapsulation of a slurry or solution of the compounds of this invention with an acceptable inert oil such as vegetable oil or light liquid petrolatum.

Pharmaceutically acceptable formulations of the disodium salts of the compounds of the present invention include: soft elastic capsules (SEC) containing a suspension of the salt; salt tablets; salt spray coated sucrose beads; or salt spray dried matrix with an enteric or non-enteric polymer

Formulations of the compounds of the present invention, which present the compounds in free acid form, preferably contain the free acid in non-crystalline form. Examples of such formulations include: soft elastic capsules containing free acid solution; non-crystalline spray dried matrix of the free acid with an enteric or non-enteric polymer; or a solid non-crystalline matrix of free acid in polyethyleneglycol (PEG) or Gelucire 44/14 (Gattefosse, Saint Priest, France).

Syrups are prepared by dissolving the compounds of this invention in an aqueous vehicle and adding sugar, aromatic flavoring agents and preservatives. Elixirs are prepared using a hydroalcoholic vehicle such as ethanol, suitable sweeteners such as sugar or saccharin and an aromatic flavoring agent. Suspensions are prepared with an aqueous vehicle and a suspending agent such as acacia, tragacanth, or methyl cellulose.

When the compounds of this invention are administered parenterally, they can be given by injection or by intravenous infusion. An effective amount is from about 0.1 mg to 100 mg per kg of body weight per day. Parenteral solutions are prepared by dissolving the compounds of this invention in liquid vehicle and filter sterilizing the solution before placing in a suitable sealable vial or ampule. Parenteral suspensions are prepared in substantially the same way except a sterile suspension vehicle is used and the compounds of this invention are sterilized with ethylene oxide or suitable gas before it is suspended in the vehicle.

The exact route of administration, dose, or frequency of administration would be readily determined by those skilled in the art and is dependent on the age, weight, general physical condition, or other clinical symptoms specific to the patient to be treated.

Patients to be treated would be those individuals: 1) infected with one or more than one strain of a human immunodeficiency virus as determined by the presence of either measurable viral antibody or antigen in the serum and 2) having either an asymptomatic HIV infection or a symptomatic AIDS defining infection such as i) disseminated histoplasmosis, ii) isoporiasis, iii) bronchial and pulmonary-candidiasis including pneumocystis pneumonia, iv) non-Hodgkin's lymphoma, or v) Kaposi's sarcoma and being less than sixty years old; or having an absolute CD4+lymphocyte count of less than 500/mm³ in the peripheral blood. Treatment would consist of maintaining an inhibitory level of the compounds of this invention in the patient at all times and would continue until the occurrence of a second

symptomatic AIDS defining infection indicates alternate therapy is needed.

The utility of representative compounds of the present invention has been demonstrated in the biological tests described below:

The HIV protease screening assay is based on fluorescently labeled substrate which can be resolved from nonlabeled cleavage product using special beads coated with streptavidin. The substrate is biotinylated at the amino terminal arginine and fluorescently labeled with fluorescein isothiocynate (FITC) at the carboxyl terminal lysine. This assay has been employed to detect novel, nonpeptidic inhibitors of HIV-1 protease. Substrate (20  $\mu$ l of 0.2  $\mu$ M), sample (10  $\mu$ l of desired concentration), and enzyme (10  $\mu$ l of 0.1  $\mu$ M) are added to a 96 well pandex plate. The assay is run <sup>15</sup> in 0.1M sodium acetate buffer at pH 5.5 in the presence of 1.0M sodium chloride and 0.05% NP-40 with incubated in the dark for one hour at room temperature. Streptavidin coated polystyrene beads  $\{40 \mu l \text{ of } 0.1\% \text{ (w/v)}\}$  are added and the plate is incubated in the dark for an additional half 20 hour. The labeled cleavage product is separated from the unreacted substrate via filtration and is read on the Idexx screen machine. The data are analyzed by appropriate computer algorithms to ascertain percent inhibition values.

Determination of  $K_i$  values utilizes the same materials and equipment employed for percent inhibition studies. Two-fold serial dilutions are made for a given inhibitor from 2, 3 or 4 starting concentrations with a total of 24, 36 or 48 individual inhibitor concentrations. These dilutions are performed utilizing the BioMek robotics system. The assay consists of 10  $\mu$ L of 40 nM HIV-1 protease, 10  $\mu$ L of the various inhibitor concentrations, and 20  $\mu$ L of 200  $\mu$ M substrate (40 uL total). The reaction is allowed to proceed for 90 min at room temperature, terminated with 40  $\mu$ L of avidin beads and processed (supra vide). An inhibitor with a known  $K_i$  is run in parallel to verify the validity of the assay. The data is processed utilizing a computer program employing a nonlinear least square analysis of the data to generate the  $K_i$  values.

The % inhibition values and/or K<sub>i</sub> values of representative compounds of the present invention tested in the HIV protease screening assay are listed in Table I below.

In the enzyme inhibition assay described above, the sensitivity of  $K_i$  value determination is in part limited by the 45 ability to continue to lower the enzyme concentration for compounds with high binding affinity. To prevent de-dimerization at low enzyme concentration, a tandemly linked enzyme is prepared in which the two monomers are covalently linked by an appropriate stretch of amino acid 50 residues. Using the latter enzyme, the sensitivity of the inhibition assay is improved since much lower enzyme concentration can be utilized, as compared to the condition using the wild-type enzyme.

Protocol for  $K_i$  value determination with tandem HIV 55 protease: Due to the greater stability (no dedimerization) of the single chain tethered (tandem) HIV protease enzyme, in which the two monomeric units are engineered to be linked by a polypeptide stretch, the method for the determination of  $K_i$  values for inhibitors uses very low concentrations of enzyme (0.2 nM) and increased incubation times (96 hours) at room temperature to improve the sensitivity in the measurement of  $K_i$  values for very potent inhibitors. The starting inhibitor concentrations are determined based on preliminary enzyme inhibition screening results which estimate the expected potency of the inhibitor. Inhibitor concentrations are then prepared using the Biomek 1000 (Beckman) and the

Quadra 96 (Tomtec). Substrate (biotinylated at the amino terminal arginine and fluorescently labeled with fluorescein at the carboxyl terminal lysine), inhibitor and the tandem enzyme are allowed to react in solution at pH 5.5 (buffers identical to those used with the native dimeric enzyme) in the dark for 96 hours. Streptavidin coated polystyrene beads are added to stop the reaction. The labeled cleavage product is separated from unreacted substrate via filtration. Residual fluorescence is quantitated with the Idexx SM2000 (Idexx) and the resulting data are analyzed using the NLLSF program.

The % inhibition values and/or  $K_i$  values of representative compounds of the present invention tested in the HIV protease screening assay and/or tandem HIV protease assay are listed in Table II below.

Several compounds of the present invention, such as N-[3-(1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide were tested in known human cell lines, such as human T-cell lines, e.g., MT4 and H9, which were infected with HIV-1<sub>IIB</sub>, and certain of these compounds were further tested in peripheral blood mononuclear cells (PBMC), which were infected with HIV-1<sub>JRCSF</sub> (a clinical isolate). The compounds were found to inhibit retroviral replication.

The following compounds of the present invention are preferred:

- 5-Cyano-N-[3-(1-[5,6-dihydro-4-hydoxy-2-oxo-6-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-phenethyl-6propyl-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide
- N-[3-(1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran- 3-yl)-2,2-dimethylpropyl)phenyl]-5-cyanopyridine-2-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide

- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-5hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethyl-propyl) phenyl]-5-cyano-2-pyridinesulfonamide
- N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide
- N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethyl-propyl) phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- 5-cyano-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- 5-amino-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl) -2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-[6-(R or S)-propyl]-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide, or (3R or S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-35 pyridinesulfonamide
- 5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-phenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide, or (3R or S, 6R or S)-N-[3-[1-(5, 6-Dihydro-4-hydroxy-2-oxo-6-propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide
- 5-Amino-N-[3-(1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-55 phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2, 2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3-(1-[5,6-dihydro-6,6-diisobutyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- N-[3(R or S)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(R or S)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2-65 dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,

- 5-Cyano-N-[3-(1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl] cyclopropylmethyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2, 2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl] propyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R or S)-(1-[6(R or S)-(2-[4-fluorophenyl] ethyl)-5,6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- N-[3(R or S)-(1-[5,6-Dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- 5-Amino-N-[3(R or S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(R or S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) phenyl]-2-pyridinesulfonamide,
- N-[3(R or S)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3(R or S)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide,
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide.
- N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis (2-phenylethyl)-2H-pyran-3-yl)-2,2dimethylpropyl}phenyl]-5-aminopyridine-2sulfonamide
- N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis (2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis (2-phenylethyl)-2H-pyran-3-yl)-2,2dimethylpropyl}phenyl]-5-cyanopyridine-2sulfonamide
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide

The following compounds of the present invention are more preferred:

- 5-Cyano-N-[3-(1-[5,6-dihydro-4-hydoxy-2-oxo-6-(2phenethyl)-6-propyl-2H-pyran-3-yl]-2,2dimethylpropyl)phenyl]-2-pyridinesulfonamide
- 5-cyano-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2Hpyran-3-yl]-2,2-dimethylpropyl]phenyl]-2pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]phenyl]-2-pyridinesulfonamide, or (3R or S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6-dipropyl-2Hpyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2- 15 pyridinesulfonamide
- 5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-npropyl-2H-pyran-3-yl]-propyl]-phenyl]-2pyridinesulfonamide, or (3R or S, 6R or S)-N-[3-[1-(5, 6-Dihydro-4-hydroxy-2-oxo-6-propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2pyridinesulfonamide
- 5-Cyano-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo- <sup>25</sup> 6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2, 2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- N-3(R or S)-1-(5,6-Dihydro-4-hydroxy-2-oxo-6(R or S)-1-(5,6-Dihydro-4-hydro-4-hydroxy-2-oxo-6(R or S)-1-(5,6-Dihydro-4-hydro-4-hydroxy-2-oxo-6(S)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4sulfonamide,
- $N-3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-2$ R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- $N-3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } S)-$ S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2sulfonamide,
- $N-\{3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } ^{40})\}\}$ S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide, and
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-45 dipropyl-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide.

The following compounds of the present invention are most preferred (see Chart EEE):

- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or 50 mL is milliliter. R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide of formula EEE-1,
- $N-[3-{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dihydro-2$ dipropyl-2H-pyran-3-yl)propyl}phenyl]-5 cyanopyridine-2-sulfonamide of formula EEE-2,
- N-[3(R or S)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(R or S)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4sulfonamide of formula EEE-3,
- 5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-npropyl-2H-pyran-3-yl]-propyl]-phenyl]-2pyridinesulfonamide of formula EEE-4, or (3R or S, 6R or S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide

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5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]phenyl]-2-pyridinesulfonamide of formula EEE-5, or (3R or S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6dipropyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide.

Also the following compounds of the present invention, which are readily prepared by the synthetic procedures set out herein, are most preferred:

- (3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6dipropyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide
- (3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-propyl-6phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide.

### DESCRIPTION OF THE PREFERRED **EMBODIMENTS**

In the Preparations and Examples below and throughout 20 this document:

°C. is degrees Centigrade.

<sup>1</sup>H-NMR is proton nuclear magnetic resonance spectrum. <sup>13</sup>C-NMR is carbon nuclear magnetic resonance spectrum. δ is chemical shift (parts per million) relative to TMS.

AlCl<sub>3</sub> is aluminum chloride.

Anal. is analytical data.

Br is benzyl.

CBZ is benzyloxycarbonyl.

CDCl<sub>3</sub> is deuterio-chloroform.

CD<sub>3</sub>OD is deuterio-methanol.

CH<sub>2</sub>Cl<sub>2</sub> is methylene chloride. cm<sup>-1</sup> is reciprocal centimeters.

CuBr<sub>2</sub> is cupric bromide.

DMSO is dimethylsulfoxide.

35 DMSO<sub>D6</sub> is deuterio dimethylsulfoxide.

EI MS is electron impact mass spectroscopy.

EtOAc is ethyl acetate.

Et<sub>3</sub>Al is triethyl aluminum.

FAB MS is fast-atom-bombardment mass spectroscopy.

HCl is hydrochloric acid.

H<sub>2</sub>O is water.

HOBT is 1-hydroxybenzotriazole hydrate.

HRMS is high-resolution mass spectroscopy.

KOH is potassium hydroxide.

M is molar (concentration).

MeOH is methanol.

Me<sub>2</sub>S is dimethyl sulfide.

mg is milligram.

MgSO<sub>4</sub> is magnesium sulfate.

mmHg is millimeter of mercury.

MP is melting point.

N is normal (concentration).

NaCl is sodium chloride.

NaOH is sodium hydroxide.

NaH is sodium hydride.

NaHCO<sub>3</sub> is sodium bicarbonate. Na<sub>2</sub>CO<sub>3</sub> is sodium carbonate.

Na<sub>2</sub>SO<sub>4</sub> is sodium sulfate.

NH<sub>4</sub>Cl is ammonium chloride.

Pd/C is palladium on charcoal.

R<sub>c</sub> is chromatographic movement relative to solvent front.

TFA is trifluoroacetic acid.

THF is tetrahydrofuran.

TMS is tetramethyl silane.

The following Preparations and Examples illustrate the present invention:

### PREPARATION 1

Cyclopropyl-(3-nitrophenyl)methanone (Formula A-2) Refer to Chart A

A 500-mL, three-necked, round-bottomed flask with a gas outlet and a 250-mL pressure-equalizing addition funnel is charged with cyclopropyl phenyl ketone of formula A-1 (30 mL) and cooled to -40° C. The addition funnel is charged with nitric acid (180 mL), which is added to the reaction mixture dropwise over 2 h. The reaction mixture is stirred another 3.5 h at -40°-0° C., and then quenched by pouring onto 500 mL of ice. The mixture is extracted with three 150-mL portions of ethyl acetate. The organic layers are combined, washed with two 250-mL portions of saturated sodium bicarbonate, dried over magnesium sulfate, filtered, and concentrated to give 41.117 g of yellow solid in an 15 orange oil. Recrystallization from 65 mL of methanol yields 20.664 g of the title product as light yellow crystals.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ8.85, 8.43, 8.33, 7.70, 2.72, 1.36–1.31, 1.20–1.14 ppm.

#### PREPARATION 2

Cyclopropyl-(3-aminophenyl)methanone (Formula A-3) Refer to Chart A

A 500-mL Parr hydrogenation flask is charged with 2.1 g of 10% platinum on carbon and a solution of the title product of Preparation 1 (20.6 g) in 250 mL of methanol. The reaction mixture is shaken for 50 min under 44 psi of hydrogen, then filtered through Celite twice. The light green solution is then concentrated to give 15.744 g of the title product as a green oil.

(Formula A-7) Refer to Chart A A 250-mL, three-necked, rountrogen inlet and a 125-mL product is charged with disoprop mL of tetrahydrofuran. The addit 4-hydroxy-2H-cycloocta[b]pyra product as a green oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.42, 7.30–7.23, 6.88, 3.83, 2.63, 1.24–1.19, 1.05–0.99 ppm.

### PREPARATION 3

N-[3-cyclopropylmethanone]benzenesulfonamide (Formula A-4) Refer to Chart A

A 500-mL, three-necked, round-bottomed flask with a nitrogen inlet is charged with the title product of Preparation 2 (15.7 g) and 200 mL of methylene chloride. Benzene-sulfonyl chloride (12 mL) and pyridine (7.8 mL) are added, and the reaction mixture is stirred at room temperature for 45 min. 10% HCl (200 mL) is added to quench the reaction. The organic layer is separated, dried over magnesium sulfate, filtered, and concentrated to give 28.638 g of orange solid. Recrystallization from 75 mL of hot methylene chloride yields the title product (22.264 g) as a pink solid.

Physical characteristics are as follows:

MP 98°-101° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.81–7.73, 7.62, 7.55–7.35, 2.60, <sup>50</sup> 1.30–1.25, 1.10–1.03 ppm,

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ200.4, 138.8, 137.2, 133.0, 129.5, 129.0, 127.0, 125.1, 124.7, 120.5, 17.3, 12.1 ppm.

IR (mineral oil) 3239, 3222, 1653, 1449, 1339, 1259, 1176, 1165, 1093, 939, 687 cm<sup>-1</sup>.

Elemental analysis, found: C, 63.70; H, 5.01; N, 4.78.

MS (EI) m/e 301, 260, 160, 141, 77.

For high resolution, found: 301.0772.

### PREPARATION 4

N-[3-cyclopropylmethanol]benzenesulfonamide (Formula A-5) Refer to Chart A

A 500-mL, three-necked, round-bottomed flask with a nitrogen inlet is charged with the title compound of Preparation 3 (21.133 g), 200 mL of tetrahydrofuran, and 100 mL 65 of ethanol. The flask is cooled to 0° C. in an ice bath, and sodium borohydride (10.6 g) is added in small portions over

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20 minutes. The reaction mixture is stirred at room temperature for ca. 18 h, and then cooled again in an ice bath to 0° C. 10% HCl (100 mL) is added dropwise over 45 min, and the mixture is stirred another 1 h at 0° C. The reaction mixture is then extracted with three 100-mL portions of methylene chloride. The organic layers are combined, dried over magnesium sulfate, filtered and concentrated to give 25.015 g of pale yellow oil. Column chromatography on 150 g of silica gel (elution with 50-65% ether in hexane followed by 2-5% methanol in methylene chloride) yields 18.692 g of the title product as a white solid.

Physical characteristics are as follows:

MP 112°-114° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.69, 7.42, 7.32, 7.25, 7.12, 7.05–6.96, 3.82, 3.19, 1.03–0.94, 0.51–0.46, 0.39–0.29, 0.19–0.16 ppm.

<sup>13</sup>C NMR (DMSO) 8147.0, 139.7, 137.4, 132.9, 129.3, 128.6, 126.8, 121.8, 118.5, 117.8, 75.0, 19.2, 3.1, 2.3 ppm.

IR (mineral oil) 3523, 3249, 1449, 732 cm<sup>-1</sup>.

Elemental analysis, found: C, 63.41; H, 5.79; N, 4.86. MS (EI) m/e 303, 275, 262, 77.

For high resolution, found: 303.0935.

### PREPARATION 5

4-Hydroxy-10-propyl-2H-cycloocta[b]pyran-2-one (Formula A-7) Refer to Chart A

A 250-mL, three-necked, round-bottomed flask with a nitrogen inlet and a 125-mL pressure-equalizing addition funnel is charged with diisopropyl amine (3.6 mL) and 15 mL of tetrahydrofuran. The addition funnel is charged with 4-hydroxy-2H-cycloocta[b]pyran-2-one of formula A-6 (2.292 g) and 35 mL of tetrahydrofuran. The flask is cooled to 0° C. in an ice bath, n-butyllithium (16.3 mL of 1.6M solution in hexanes) is added dropwise over 3 min, and the reaction mixture is stirred another 15 min at 0° C. The solution of 4-hydroxy-2H-cycloocta[b]pyran-2-one in THF is added dropwise over 35 min, and the reaction mixture is stirred for another 25 min at 0° C. Hexamethylphosphoramide (4 mL) is added in one portion, and iodopropane (1.3 mL) is added dropwise over 2 min. The reaction mixture is allowed to warm to room temperature and stirred for ca. 18 h. 30 mL of 10% HCl is added and the aqueous layer is separated. The pH of the aqueous layer is lowered from 10 to 2 with concentrated HCl, and the aqueous layer is extracted with two 50-mL portions of methylene chloride. The organic layers are combined, dried over magnesium sulfate, filtered, and concentrated to give an orange oil, which is partitioned between 100 mL of 1N sodium hydroxide and 50 mL of ether. The aqueous layer pH is adjusted from 14 to 1 with concentrated hydrochloric acid, and is then extracted with two 50-mL portions of methylene chloride. The organic layers are then combined, dried over magnesium sulfate, and concentrated to give an orange oil, which is diluted with 100 mL of ether and washed with three 25-mL portions of 10% HCl. The organic layer is then dried over magnesium sulfate, filtered, and concentrated to give 1.829 g of orange solid. Column chromatography on 100 g of silica gel (elution with 0-10% methanol in methylene chloride) gives 1.358 g of a pale orange solid. An additional column chromatography on 150 g of silica gel (elution with 10% ether and 1% acetic acid in methylene chloride) gives 0.705 g of the title product as a yellow solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ11.38, 5.68, 3.02–2.93, 2.20, 1.98–1.82, 1.73–1.58, 1.46–1.25, 1.24–1.08, 0.89 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ172.3, 168.3, 165.3, 114.8, 89.7, 38.6, 36.0, 33.3, 30.1, 27.2, 25.5, 22.9, 21.0, 13.9 ppm. IR (mineral oil) 1679, 1641, 1617, 1492 cm<sup>-1</sup>.

Elemental analysis, found: C, 70.90; H, 8.36. MS (EI) m/e 236, 208, 166. For high resolution, found: 236.1414.

### **EXAMPLE 1**

N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-10-propyl-2H-cycloocta[b]pyran-3-yl)methyl]phenyl] benzenesulfonamide (Formula A-8) Refer to Chart A

A 100-mL, three-necked, round-bottomed flask with a 35-mL pressure-equalizing addition funnel filled with 3 A 10 molecular sieves and fitted with a reflux condenser and a nitrogen inlet is charged with the title compound of Preparation 5 (0.196 g), p-toluenesulfonic acid (0.040 g), and 30 mL of methylene chloride. The title product of Preparation 4 (0.252 g) is added, and the reaction mixture is heated to 15 reflux for 2 h, then stirred at room temperature for an additional hour. The reaction mixture is then diluted with 20 mL of methylene chloride and washed with 60 mL of 1:1 saturated sodium bicarbonate and brine, 30 mL of water, and 30 mL of brine. The aqueous layers are combined and  $_{20}$ extracted with 30 mL of methylene chloride. The organic layers are then combined, dried over magnesium sulfate, filtered, and concentrated to give 0.576 g of crude material. Column chromatography on 35 g of silica gel (elution with 20-80% ether in hexane) yields 0.096 g of the title com- 25 pound as a white solid.

Physical characteristics are as follows: MP 87°-90° C. (decomposition). MS (EI) m/e 521, 493, 380, 275, 262, 249, 144, 77. For high resolution, found: 521.2236.

#### EXAMPLES 2-7

Following procedures analogous to those described above, the following additional compounds of the present invention are prepared:

- 2) 4-Cyano-N-[3-(R or S)-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-(R or S)-10-propyl-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]benzenesulfonamide
- 3) 4-Cyano-N-[3-(R or S)-[cyclopropyl(5,6,7,8,9,10-hexahydro-(R or S)-10-cyclopropylmethyl-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]benzenesulfonamide
- 4) 4-Cyano-N-[3-(R or S)-[cyclopropyl(5,6,7,8,9,10-hexahydro-(R or S)-10-benzyl-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl] benzenesulfonamide
- 5) N-[3-(R or S)-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-(R or S)-10-propyl-2H-cycloocta[b] 50 pyran-3-yl)methyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide
- 6) N-[3-(R or S)-[cyclopropyl(5,6,7,8,9,10-hexahydro-(R or S)-10-cyclopropylmethyl-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]-1-methyl-1H- 55 imidazole-4-sulfonamide
- N-[3-(R or S)-[cyclopropyl(5,6,7,8,9,10-hexahydro-(R or S)-10-benzyl-4-hydroxy-2-oxo-2H-cycloocta[b] pyran-3-yl)methyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide

### PREPARATION 6

(3-Benzaldehyde)-carbamic acid, phenylmethyl ester (Formula B-2) Refer to Chart B

A flask with a nitrogen inlet is charged with sodium 65 bicarbonate (10.4 g) in 200 mL of THF and 200 mL of water, and m-aminobenzaldehyde of formula B-1 (10.0 g) and

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benzyl chloroformate (13.6 mL) are added sequentially. The mixture is stirred at room temperature for 40 min. Ether is then added, and the organic layer is separated, washed with saturated sodium bicarbonate, dried over sodium sulfate, filtered and concentrated to give a brown oil. Column chromatography on 300 g of silica gel yields 16.3 g of the title compound as a pale yellow oil. An analytical sample is crystallized from ethyl acetate-hexane.

Physical characteristics are as follows:

MP 100°-104° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ9.98, 7.91, 7.69, 7.59, 7.43–7.35, 6.83, 5.23 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ191.8, 153.0, 138.6, 137.1, 135.6, 129.7, 128.6, 128.4, 128.3, 124.6, 124.2, 119.1, 67.2 ppm. IR (mineral oil) 3269, 2954, 2925, 2868, 2855, 1729, 1682, 1597, 1560, 1465, 1455, 1326, 1294, 1237, 1229, 1170, 1155, 1048, 695 cm-1.

Elemental analysis, found: C, 70.74; H, 5.14; N, 5.33. MS (EI) m/e 255, 211, 91.

For high resolution, found 255.0900.

### PREPARATION 7

[3-(1-Hydroxy-3-methylbutyl)phenyl]-carbamic acid, phenylmethyl ester (Formula B-3 wherein R<sub>1</sub> is isobutyl) Refer to Chart B

A flask with a nitrogen inlet is charged with the title compound of Preparation 6 (4.0 g) and 60 mL of dry tetrahydrofuran. The mixture is cooled to 0° C., and isobutyl magnesium chloride (17.2 mL) is added. The reaction mixture is then allowed to warm to room temperature and stir for 2 hours. Saturated ammonium chloride is added to quench the reaction, and the mixture is partitioned between ether and water. The organic layer is washed with water and concentrated to give 5.78 g of pale yellow oil. The crude material is crystallized from ethyl acetate-hexane to yield 4.13 g of the title compound as white crystals.

Physical characteristics are as follows:

MP 73°-77° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.41–7.33, 7.25, 7.05, 6.74, 5.19, 4.73–4.65, 1.91, 1.73–1.65, 1.47, 0.93 ppm.

IR (Nujol) 3400, 3249, 3085, 2953, 2925, 2869, 2855, 1697, 1615, 1602, 1563, 1450, 1283, 1245, 1177, 1067, 1017, 798, 773, 740, 696 cm<sup>-1</sup>.

Elemental analysis, found: C, 72.58; H, 7.25; N, 4.55. MS (El) m/z 313, 257, 213, 91.

## PREPARATION 8

[3-[1-(5,6,7,8,9,10-Hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)-3-methylbutyl]phenyl-carbamic acid, phenylmethyl ester (Formula B-5 wherein  $R_1$  is isobutyl) Refer to Chart B

A 200-mL, three-necked flask with a Dien-Stark trap and a nitrogen inlet is charged with p-toluenesulfonic acid (0.66 g) and toluene (100 mL) and warmed to reflux to collect 20 mL in the Dien-Stark trap. The reaction mixture is cooled to room temperature, and the trap is emptied. 4-Hydroxy-2H-cycloocta[b]pyran-2-one of formula B-4 (2.48 g) and the title compound of Preparation 7 (4.0 g) are added to the reaction mixture and then heated to reflux for 6.5 h. The reaction mixture is allowed to stand at room temperature overnight, then poured into 350 mL of ethyl acetate, washed with two 25-mL portions of water, 25 mL of saturated sodium bicarbonate, and 25 mL of water. The organic layer is concentrated to give 7.9 g of yellow oil. Column chromatography on 150 g of silica gel (elution with 10-50%

ethyl acetate in hexane) gives 0.217 g of the title product as an off-white foam.

Physical characteristics are as follows:

MP 73°-78° C. (decomposition).

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.38–7.25, 7.13, 6.72, 6.01, 5.19, 5 4.48, 2.58, 2.41, 1.93, 1.74, 1.62-1.33, 0.96 ppm.

#### PREPARATION 9

(R or S)-[3-[1-(5,6,7,8,9,10-Hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)-3-methylbutyl]phenyl]- 10 carbamic acid, phenylmethyl ester (Formula B-5 wherein R<sub>1</sub> is isobutyl) Refer to Chart B

A stock solution of the title compound of Preparation 8 (32 mg/mL) in 30% isopropyl alcohol and 0.1% acetic acid in hexane is chromatographed on a 2.0×25 cm (R, R) 15 Whelk-O 1 column at 2 mL per injection using an automated chromatographic system. The eluant is monitored at 310 nm, the flow rate was 10 mL/min and appropriate fractions from multiple injections combined and concentrated in vacuo to give snowy white solids.

Physical characteristics are as follows:

The retention time of the title compound is 18.8 min.

### PREPARATION 10

(R or S)-[3-[1-(5,6,7,8,9,10-Hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)-3-methylbutyl]phenyl]carbamic acid, phenylmethyl ester (Formula B-5 wherein R, is isobutyl) Refer to Chart B

The title compound of Preparation 8 is separated as described in Preparation 9 above.

Physical characteristics are as follows:

The retention time of the title compound is 22.1 min.

### PREPARATION 11

(R or S)-3-[1-(3-Aminophenyl)-3-methylbutyl]-5,6,7,8,9, 10-hexahydro-4-hydroxy-2H-cycloocta[b]pyranone 35 (Formula B-6 wherein R<sub>1</sub> is isobutyl) Refer to Chart B

A flask with a nitrogen inlet is charged with a solution of the title compound of Preparation 9 (0.637 g) in 6 mL of ethanol. Cyclohexene (6 mL) and 10% palladium on carbon (0.16 g) are added, and the reaction mixture is heated at 40 reflux for 2 h. The mixture is then filtered through Celite and concentrated to give 0.205 g of the title compound as an off-white foam.

Physical characteristics are as follows:

MP 158°-162° C.

MS (EI) m/z 355, 312, 299, 161, 106

For high resolution, found: 355.2144.

#### **EXAMPLE 8**

(R or S)-N-[3-1-(5,6,7,8,9,10-Hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)-3-methylbutyl]phenyl]1methyl-1H-imidazole-4-sulfonamide (Formula B-7 wherein R<sub>1</sub> is isobutyl and R<sub>2</sub> is 1-methylimidazole) Refer to Chart

A flask with a nitrogen inlet is charged with the title compound of Preparation 11 (0.095 g), 1-methylimidazole- 55 4-sulfonyl chloride (0.048 g), and 5 mL of methylene chloride (CH<sub>2</sub>Cl<sub>2</sub>). Pyridine (0.53 mL) is added, and the reaction mixture is stirred at room temperature for ca. 18 h. A precipitate forms, which is filtered to give 0.097 g of a white solid. Recrystallization from methanol-chloroform 60 yields 0.065 g of the title compound as a white powder.

Physical characteristics are as follows:

MP 207°-210° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ10.4, 10.0, 7.70, 7.11, 7.05, 6.92, 4.21, 3.64, 2.54, 2.16, 1.62, 1.53, 1.43, 1.34, 0.85 ppm.

MS (EI) m/z 499, 456, 443, 306, 251, 160, 145

For high resolution, found: 499.2151

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#### PREPARATION 12

(R or S)-3-[1-(3-Aminophenyl)-3-methylbutyl]-5,6,7,8,9, 10-hexahydro-4-hydroxy-2H-cycloocta[b]pyranone (Formula B-6 wherein R<sub>1</sub> is isobutyl) Refer to Chart B

Following the general procedure of Preparation 11, and making non-critical variations, but substituting the title product of Preparation 10 for the title product of Preparation 9, 0.189 g of the title compound is obtained as a grey solid.

Physical characteristics are as follows:

MS (EI) m/z 355, 312, 299, 161

For high resolution, found: 355.2135

#### EXAMPLE 9

(R or S)-N-[3-1-(5,6,7,8,9,10-Hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)-3-methylbutyl]phenyl]-1methyl-1H-imidazole-4-sulfonamide (Formula B-7 wherein R<sub>1</sub> is isobutyl and R<sub>2</sub> is 1-methylimidazole) Refer to Chart

Following the general procedure of Example 8, and making non-critical variations, but substituting the title product of Preparation 12 for the title product of Preparation 11, 0.047 g of the title compound is obtained as a white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ10.45, 10.06, 7.70, 7.11, 7.05, 6.94, 25 4.21, 3.64, 2.55, 2.16, 1.62, 1.53, 1.42, 1.35, 0.86 ppm.

MS (EI) m/z 499, 456, 443, 354, 306, 160, 145 For high resolution, found: 499.2146

### PREPARATION 13

[3-(Cyclopropyl-hydroxymethyl)-phenyl]-methanol (Formula C-2) Refer to Chart C

To a solution of 6.5 mL of 3-bromobenzylalcohol of formula C-1 in 900 mL of tetrahydrofuran under nitrogen at -78° C. is added 46 mL of a 1.4M solution of methyllithium in diethyl ether. The solution is stirred for 20 min and then 66 mL of a 1.6M solution of n-butyllithium in hexane is added. The solution is stirred 25 min and then 6 mL of cyclopropanecarboxaldehyde is added. The solution is stirred 1.5 h, warmed to 0° C. and stirred for 40 min. Next the solution is warmed to room temperature and stirred for 30 min. Finally the solution is heated at reflux for 1 h. The solution is poured onto 800 mL of water and acidified with concentrated HCl followed by 5% aqueous HCl to adjust the pH to approximately 6. The layers are separated and the aqueous extracted with two portions of ethyl acetate. The 45 combined organics are dried (Na<sub>2</sub>SO<sub>4</sub>) and concentrated to afford a yellow oil which is chromatographed over 900 g 230-400 mesh silica gel (2:1 ethyl acetate:hexane) to afford a 6.61 g (68%) of the desired alcohol as a yellow oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.41–7.26, 4.67, 3.99–3.96, 2.18, 1.28-1.14, 0.68

### PREPARATION 14

3-[cyclopropyl [3-[hydroxymethyl]phenyl]methyl]-5,6,7,8, 9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one (Formula C-3) Refer to Chart C

To a solution of 501 mg of the title product of Preparation 13 in 50 mL of dichloromethane in the presence of molecular sieves 3A under nitrogen is added 492 mg of 4-hydroxy-5,6,7,8,9,10-hexahydrocycloocta[b]pyran-2-one followed by 49 mg of p-toluenesulfonic acid monohydrate. The solution is heated at reflux for 2 h and then an additional 105 mg p-toluenesulfonic acid monohydrate is added and heating continued for a further hour. The solution is concentrated in vacuo to afford a white foam which is treated with water and then 1N KOH and extracted with one portion of ethyl acetate. The organic layer is washed with one portion of 1N

KOH. The combined aqueous layers are acidified with 5% aqueous HCl and extracted with three portions of ethyl acetate. The combined organics are dried (Na<sub>2</sub>SO<sub>4</sub>) and concentrated in vacuo to afford a yellow oil which is chromatographed over 180 g of 230–400 mesh silica gel (2:1 5 ethyl acetate:hexane) to afford 436 mg of the desired benzyl alcohol as a white foam.

Physical characteristics are as follows:

MP 65°-70° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.25–7.03, 4.36, 3.70–3.67, 10 2.41–2.37, 2.24–2.23, 1.53–1.50, 1.35–1.05, 0.54–0.43, 0.42–0.21, 0.07–0.02.

### PREPARATION 15

3-[Cyclopropyl [3-[bromomethyl]phenyl]methyl]-5,6,7,8,9, 10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one; and 3-[cyclopropyl[3-[chloromethyl]phenyl]methyl]-5,6,7,8,9, 10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one (Formulas C-4,5) Refer to Chart C

To a solution of 1.01 g of the title product of Preparation 14 in 70 mL of dichloromethane under nitrogen at 0° C. is added 2.00 g of triphenylphosphine and 2.58 g of carbon tetrabromide in sequence. The solution is stirred 1 h and then poured onto brine. The layers are separated and the aqueous extracted with three portions of ethyl acetate. The combined organics are dried (Na<sub>2</sub>SO<sub>4</sub>) and concentrated to afford a yellow oil which is triturated with ether. The solid is filtered off and the filtrate concentrated and chromatographed over 180 g of 230–400 mesh silica gel (1:1 hexane:ethyl acetate) to afford 374 mg of the desired title product as a mixture of bromide and chloride. The solids isolated from the filtration are chromatographed as above to afford an additional 699 mg of the title product as a mixture of bromide and chloride.

Physical characteristics are as follows:

Mass Spectrum m/e 418, 416 (M<sup>+</sup> for Br), 388, 374, 372 (M<sup>+</sup> for Cl), 337, 246, 233, 220, 207, 195, 179, 153, 143, 129.

### PREPARATION 16

3-[Cyclopropyl[3-[(phenylthio)methyl]phenyl]methyl]-5,6, 7,8,9,10-hexahydro-4-hydroxy-2H-Cycloocta[b]pyran-2-one (Formula C-6) Refer to Chart C

To a solution of 138 mg of the title products of Preparation 15 in 5 mL of dichloromethane is added 0.04 mL of thiolphenol and 0.17 mL of diisopropylethylamine in sequence. The solution is heated at reflux for 1 h and then allowed to stand at room temperature overnight. The solution is poured onto brine and treated with 5% aqueous hydrochloric acid. The layers are separated and the aqueous extracted with three portions of ethyl acetate. The combined organics are dried (Na<sub>2</sub>SO<sub>4</sub>) and concentrated to afford a yellow oil which is chromatographed over 80 g of 230–400 mesh silica gel (2:1 hexane:ethyl acetate) to afford 111 mg of the desired sulfide as a white foam.

Physical characteristics are as follows:

MP 137°-139° C.

Mass Spectrum m/e 446 (M\*), 418, 337, 295, 233, 220, 207, 185, 145, 128, 109, 91, 79, 55, 40.

# **EXAMPLE 10**

3-[Cyclopropyl[3-[(phenylsulfonyl)methyl]phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-Cycloocta[b]pyran-2-one (Formula C-7) Refer to Chart C

To a solution of 119 mg of the title product of Preparation 16 in 5 mL of tetrahydrofuran and 5 mL of methanol at 0° C. is added a solution of 279 mg of oxone in 5 mL of water. 65 The solution is stirred 2.5 h and then warmed to room temperature and stirred 2 h. The solution is filtered and the

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solids washed with chloroform. The filtrate is diluted with water and the layers are separated. The aqueous is extracted with three portions of ethyl acetate. The combined organics are dried (Na<sub>2</sub>SO<sub>4</sub>) and concentrated to afford a clear oil which is chromatographed over 80 g of 230–400 mesh silica gel (1:1 hexane:ethyl acetate) to afford 78 mg of the title product as a white foam.

Physical characteristics are as follows:

MP 80°-85° C.

Mass Spectrum m/e 479 (M<sup>+</sup>+1), 463, 450, 391, 337, 309, 207, 161, 149, 127, 115, 71, 57, 41.

Exact mass found: 479.1885.

#### EXAMPLES 11-39

The following compounds of the present invention are prepared by an analogous synthetic route to that described above:

11) 3-[cyclopropyl[3-[(4-cyanophenylsulfonyl)methyl] phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

The starting material, 4-cyanobenzenethiol, is prepared from 4-cyanobenzenesulfonyl chloride according to a general literature procedure: Wagner, A. W. Ber Deutsch Chem Ges, 99:375 (1966).

Physical characteristics are as follows:

MP 100°-105° C.

Mass Spectrum m/e 504 (M<sup>+</sup>+1), 337, 247, 207, 143.

Exact mass found 504.1843.

12) 3-[cyclopropyl[3-[(4-fluorophenylsulfonyl)methyl] phenyl]methyl]- 5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

Physical characteristics are as follows:

MP 95°-100° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.61–7.57, 7.40–7.37, 7.27–7.20, 7.13–7.07, 7.02–6.99, 6.42, 4.30, 3.88–3.85, 2.64–2.61, 2.51–2.47, 1.83–1.40, 1.40–1.27, 0.69–0.58, 0.48–0.43, 0.19–0.14.

13) 3-[cyclopropyl[3-[(4-methylphenylsulfonyl)methyl] phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

Physical characteristics are as follows:

MP 100°-105° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.37–7.34, 7.25–7.22, 7.17–7.05, 6.86–6.84, 4.15, 3.60–3.58, 2.52–2.42, 2.42–2.30, 2.28, 1.70–1.14, 0.57–0.32, 0.32–0.20, 0.06-(–)0.16.

14) 3-[cyclopropyl[3-[(4-carboxyphenylsulfonyl)methyl] phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

Physical characteristics are as follows:

MP 90°-95° C.

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Mass Spectrum m/e 523 (M\*+1), 337, 247, 207, 143. Exact mass found 523.1785.

15) 3-[cyclopropyl[3-[(2-(1-methylimidazoyl)sulfonyl) methyl]phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

Physical characteristics are as follows: MP 95°-103° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.36–7.34, 7.29–7.27, 7.14, 7.06–7.03, 6.98 (s, 1H), 6.86, 4.30, 3.73–3.70, 3.20, 2.67–2.54, 1.90–1.36, 0.71–0.50, 0.46–0.33, 0.18–0.03.

- 16) 3-[cyclopropyl[3-[(2-pyrimidinylsulfonyl)methyl] phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one
- 17) 3-[cyclopropyl[3-[(2-pyridinylsulfonyl)methyl] phenyl]methyl]-5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

- 18) 3-[cyclopropyl[3-[(1-methyl-4-imidazolylsulfonyl) methyl]phenyl]methyl]-5,6,7,8,9,10-hexahydro-4hydroxy-2H-cycloocta[b]pyran-2-one
- 19) 3-[cyclopropyl[3-[(5-cyano-2-pyridinylsulfonyl) methyl]phenyl]methyl]-5,6,7,8,9,10-hexahydro-4- 5 hydroxy-2H-cycloocta[b]pyran-2-one
- 20) 3-[cyclopropyl[3-[(2-benzimidazolylsulfonyl) methyl]phenyl]methyl]-5,6,7,8,9,10-hexahydro-4hydroxy-2H-cycloocta[b]pyran-2-one
- phenyl]methyl]- 5,6,7,8,9,10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one
- 22) 3-[cyclopropyl[3-[(4-cyanophenylsulfonyl)methyl] phenyl methyl -4-hydroxy-coumarin
- 23) 3-[cyclopropyl[3-[(2-pyridinylsulfonyl)methyl] phenyl]methyl]-4-hydroxy-coumarin
- 24) 3-[cyclopropyl[3-[(1-methyl-4-imidazolylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-coumarin
- 25) 3-[cyclopropyl[3-[(5-cyano-2-pyridinylsulfonyl) 20 methyl]phenyl]methyl]-4-hydroxy-coumarin
- 26) 3-[cyclopropyl[3-[(2-benzimidazolylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-coumarin
- 27) 3-[cyclopropyl[3-[(2-quinolinylsulfonyl)methyl] phenyl]methyl]-4-hydroxy-coumarin
- 28) 3-[cyclopropyl[3-[(4-cyanophenylsulfonyl)methyl] phenyl]methyl]-4-hydroxy-6-[1-(phenylmethyl) propyl]-2H-pyran-2-one
- 29) 3-[cyclopropyl[3-[(2-pyridinylsulfonyl)methyl] 30 phenyl]methyl]-4-hydroxy-6-[1-(phenylmethyl) propyl]-2H-pyran-2-one
- 30) 3-[cyclopropyl[3-[(1-methyl-4-imidazolylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-6-[1-(phenylmethyl)propyl]-2H-pyran-2-one
- 31) 3-[cyclopropyl[3-[(5-cyano-2-pyridinylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-6-[1-(phenylmethyl)propyl]-2H-pyran-2-one
- 32) 3-[cyclopropyl[3-[(2-benzimidazolylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-6-[1-40 (phenylmethyl)propyl]-2H-pyran-2-one
- 33) 3-[cyclopropyl[3-[(2-quinolinylsulfonyl)methyl] phenyl]methyl]-4-hydroxy-6-[1-(phenylmethyl) propyl]-2H-pyran-2-one
- 34) 3-[cyclopropyl[3-[(4-cyanophenylsulfonyl)methyl] 45 phenyl]methyl]-4-hydroxy-6-(2-phenylethyl)-6-(1propyl)-5,6-dihydro-2H-pyran-2-one
- 35) 3-[cyclopropyl[3-[(2-pyridinylsulfonyl)methyl] phenyl]methyl]-4-hydroxy-6-(2-phenylethyl)-6-(1propyl)-5,6-dihydro-2H-pyran-2-one
- 36) 3-[cyclopropyl[3-[(1-methyl-4-imidazolylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-6-(2-phenylethyl)-6-(1-propyl)-5,6-dihydro-2H-pyran-2-one
- 37) 3-[cyclopropyl[3-[(5-cyano-2-pyridinylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-6-(2-phenylethyl)-6-(1-propyl)-5,6-dihydro-2H-pyran-2-one
- 38) 3-[cyclopropyl[3-[(2-benzimidazolylsulfonyl) methyl]phenyl]methyl]-4-hydroxy-6-(2-phenylethyl)-6-(1-propyl)-5,6-dihydro-2H-pyran-2-one
- 39) 3-[cyclopropyl[3-[(2-quinolinylsulfonyl)methyl] phenyl]methyl]-4-hydroxy-6-(2-phenylethyl)-6-(1propyl)-5,6-dihydro-2H-pyran-2-one

### PREPARATION 17

5,6-Dihydro-4-Hydroxy-6-phenethyl-6-propyl-2H-pyran-2- 65 one (Formula D-1: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl) Refer to Chart D

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Methyl acetoacetate (1.47 mL) is added to a suspension of sodium hydride (567 mg, 60% dispersion in mineral oil) in THF (30 mL) at 0° C. After 15 minutes, n-butyl lithium (8.5 mL, 1.6M solution in hexane) is added dropwise and the reaction is stirred 15 minutes. 1-Phenyl-3-hexanone (2.0 g) is then added via syringe all at once to the reaction mixture. The reaction is stirred an additional hour, then poured into a saturated ammonium chloride solution. It is extracted with EtOAc, dried over anhydrous sodium sulfate and evaporated 21) 3-[cyclopropyl[3-[(2-quinolinylsulfonyl)methyl] 10 in vacuo. The material obtained is dissolved in THF (25 mL) and a 0.1N sodium hydroxide (113 mL) solution is added. After stirring three hours, the mixture is extracted with ethyl acetate (1x). The aqueous layer is adjusted to pH 3 with hydrochloric acid, then extracted with CH<sub>2</sub>Cl<sub>2</sub> (3×25 mL), 15 dried over anhydrous magnesium sulfate and evaporated to afford the title product as a white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>): δ0.96, 1.21, 1.48, 1.72, 1.98, 2.73, 3.43, 7.15–7.32.

Anal. Found: C, 73.77; H, 7.96.

#### PREPARATION 18

4-Hydroxy-3-[1-(3-nitrophenyl)-propyl]-5,6-dihydro-6phenethyl-6-propyl-2H-pyran-2-one (Formula D-4: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) Refer to Chart D

To a solution of the title product of Preparation 17 (Formula D-1: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl) (1 g) and 3-nitrobenzaldehyde (Formula D-2) (581 mg) in dry THF at 0° C. is added AlCl<sub>3</sub> (1.0 g) as one solid portion. The cooling bath is removed and the yellow solution is allowed to stir at room temperature for 2 hrs. The reaction mixture is quenched by the addition of solid Na<sub>2</sub>CO<sub>3</sub>-10H<sub>2</sub>O (2.2 g) and vigorously stirred for 5 min. The mixture is filtered through celite with ether and the filtrate is evaporated to dryness in vacuo. The benzylidene intermediate of the formula D-3 and CuBr-Me<sub>2</sub>S (237 mg) are dissolved in dry THF and a solution of Et<sub>3</sub>Al (4.23 mL; 1M in hexane) is added at room temperature, dropwise over 5 min. When the reaction is complete (as determined by tlc), it is quenched by the addition of water and the reaction mixture is transferred to a separatory funnel with ether. The aqueous layer is extracted with ether (3×15 mL) and the combined organic layers are washed with brine, dried (MgSO<sub>4</sub>), filtered and evaporated in vacuo to provide an oil. Flash chromatography on silica gel with Hexanes/EtOAc (3:1) provides 1.1 g of the title product as a light yellow foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.93, 1.37, 1.74, 1.82-2.14, 2.29, 2.52-2.71, 4.19, 6.98-7.24, 7.44, 7.72, 8.02, 8.26.

# PREPARATION 19

3-[Cyclopropyl-(3-nitrophenyl)-methyl]-4-hydroxy-5,6dihydro-6-phenethyl-6-propyl-2H-pyran-2-one (Formula D-4: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is cyclopropyl) Refer to Chart D

To a solution of the title product of Preparation 17 (Formula D-1: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl) (1 g) and 3-nitrobenzaldehyde (Formula D-2) (581 mg) in dry THF at 0° C. is added AlCl<sub>3</sub> (1.0 g) as one solid portion. The cooling bath is removed and the yellow solution is allowed to stir at room temperature for 2 hrs. The reaction mixture is quenched by the addition of solid Na<sub>2</sub>CO<sub>3</sub>-10H<sub>2</sub>O (2.2 g) and vigorously stirred for 5 min. The mixture is filtered through celite with ether and the filtrate is evaporated to dryness in vacuo. The benzylidene intermediate of formula D-3 and CuBr-Me<sub>2</sub>S (237 mg) are dissolved in dry THF

and cooled to -78° C. A solution of cyclopropylmagnesium bromide (15.6 mL; 0.25M in THF) is added dropwise over 10 min and the reaction mixture is stirred for 30 min. The reaction is quenched by the addition of water and neutralized by the addition of 1N HCl. The reaction mixture is transferred to a separatory funnel with ether and the aqueous layer is extracted with ether (3×15 mL). The combined organic layers are washed with brine, dried (MgSO<sub>4</sub>), filtered and evaporated in vacuo to provide an oil. Flash chromatography on silica gel with Hexanes/EtOAc (3:1) 10 provides 0.9 g of the title product as a light yellow foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.25, 0.53, 0.74, 0.94, 1.41, 1.68–2.13, 2.57–2.72, 3.38, 7.04–7.23, 7.46, 7.82, 15 1.45–2.11, 2.43–2.68, 3.24, 3.64, 3.94, 6.72–7.51. 8.03, 8.30.

### PREPARATION 20

3-[1-(3-Aminophenyl)-propyl]-4-hydroxy-5,6-dihydro-6phenethyl-6-propyl-2H-pyran-2-one (Formula D-5: R, is phenethyl, R2 is propyl, R3 is ethyl) Refer to Chart D

To a solution of the title product of Preparation 18 (Formula D-4:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is ethyl) (350 mg) in MeOH at room temperature is added 10% Pd/C (35 mg) and ammonium formate (521 mg). The resulting 25 mixture is stirred for 2 hrs. and then filtered through celite with CH<sub>2</sub>Cl<sub>2</sub>. The filtrate is evaporated in vacuo and the residue is triturated with CH<sub>2</sub>Cl<sub>2</sub> (3×10 mL). The combined organic solution is filtered and evaporated in vacuo to provide the 325 mg of the title compound as a light yellow 30

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ0.89, 1.40, 1.64–2.07, 2.20, 2.62, 3.94, 6.54, 6.72–7.25.

The compounds of formula D-5, wherein R<sub>1</sub> is propyl, R<sub>2</sub> is propyl and R<sub>3</sub> is ethyl or t-butyl are prepared by analogous procedures.

Physical characteristics of the compound of the formula D-5, wherein R<sub>1</sub> is and R<sub>2</sub> are propyl and R<sub>3</sub> is ethyl, are as 40

<sup>1</sup>H NMR: 0.9, 1.3, 1.5–1.8, 2.0, 2.2, 2.5, 3.9, 4.5, 6.5, 6.8, 7.0 ppm

TLC R: 0.32 (10% ethyl acetate in dichloromethane).

### **EXAMPLE 40**

4-Cyano-N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]benzenesulfonamide (Formula D-6: R, is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 4-cyanophenyl) Refer to Chart D

To a solution of the title product of Preparation 20 50 (Formula D-5:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is ethyl) (30 mg) and 4-cyanobenzenesulfonyl chloride of formula D-7, wherein R<sub>4</sub> is cyanophenyl, (16.1 mg) in CH<sub>2</sub>Cl<sub>2</sub> (1 mL) at room temperature is added pyridine (13  $\mu$ L) via syringe. The resulting solution is stirred for 3 hrs, after which the starting amine is consumed. The mixture is flash chromatographed on silica gel with the 5% EtOAc in CH2Cl2 to provide 21 mg of the title product as a white foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ0.6–1.1, 1.2–2.2, 2.4–2.7, 3.86-4.01, 6.89-7.45, 7.66-7.92.

HRMS found: 559.2267.

#### **EXAMPLE 41**

N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1-methyl-1H-

imidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>2</sub> is ethyl, R<sub>4</sub> is 1-methylimidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5; R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein R<sub>4</sub> is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers and tautomerism.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) 80.75–0.96, 1.17–1.43, HRMS found: 538.2383.

### **EXAMPLE 42**

N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6dihydro-2H-pyran-3-yl)-propyl]-phenyl]-8quinolinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is quinolin-8-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 8-quinolinesulfonyl chloride of formula D-7 wherein 44 is quinolin-8-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH2Cl2.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers and tautomerism.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ0.66, 0.90, 1.17–1.44, 1.58-2.03, 2.38-2.64, 3.77, 6.68-7.27, 7.35-7.69, 8.02, <sub>35</sub> 8.26, 9.14.

HRMS found: 585.2402.

### **EXAMPLE 43**

N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2Hpyran-3-yl)-propyl]-phenyl]-8-quinolinesulfonamide (Formula D-6: R<sub>1</sub> is R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is quinolin-8-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 8-quinolinesulfonyl chloride of formula D-7 wherein R<sub>4</sub> is quinolin-8-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers and tautomerism.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.67, 0.85, 1.27, 1.54, 2.01, 3.73, 6.78, 6.90, 7.04, 7.57, 8.12, 8.29, 8.38, 9.13. HRMS found: 523.2276.

Anal. found: C, 66.09; H, 6.60; N, 5.13.

### **EXAMPLE 44**

N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2Hpyran-3-yl)-propyl]-phenyl]-1-methyl-1Himidazolesulfonamide (Formula D-6: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 1-methylimidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield

the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.88, 1.32, 1.64, 1.93, 2.16, 2.56, 3.68, 3.91, 6.87, 7.03, 7.14, 7.53, 7.64. HRMS found: 476.2223.

### **EXAMPLE 45**

4-Fluoro-N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 4-fluorophenyl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 4-fluoro-benzenesulfonyl chloride of formula D-7 wherein R<sub>4</sub> is 4-fluorophenyl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers and tautomerism.

<sup>1</sup>H NMR (300 MHz, CDCL<sub>3</sub>) δ0.51–1.03, 1.15–1.73, 1.81–2.48, 2.73, 3.91, 6.69, 6.88, 7.09, 7.78. HRMS found: 490.2085.

4-Cyano-N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 4-cyanophenyl) Refer to Chart D

**EXAMPLE 46** 

The title compound is prepared from the amine of Preparation 20 (Formula D-5: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 4-cyano-benzenesulfonyl chloride of formula D-7 wherein R<sub>4</sub> is 4-cyanophenyl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers and tautomerism.

<sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ0.68–0.96, 1.15–1.42, 1.44–1.76, 1.83–2.12, 3.18, 3.88, 6.69–7.18, 7.71, 7.85. HRMS found: 497.2126.

### **EXAMPLE 47**

N- $\{3-[1-(4-hydroxy-6,6-diisobutyl-2-oxo-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is isobutyl, R<sub>2</sub> is isobutyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 1-methylimidazol-4-yl) Refer to Chart D$ 

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is propyl,  $R_2$  is isobutyl,  $R_3$  is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after 55 flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.82–0.94, 1.52–1.83, 1.86–2.03, 2.06–2.22, 2.60, 3.68, 3.92, 6.87, 7.03, 7.16, 7.56, 7.65.

HRMS found: 504.2531.

Anal. found: C, 62.03; H, 7.43; N, 8.20.

### **EXAMPLE 48**

N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H- $_{65}$  pyran-3-yl)-cyclopropylmethyl]-phenyl]-1-methyl-1H-imidazolesulfonamide (Formula D-6:  $R_{1}$  is propyl,  $R_{2}$  is

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propyl,  $R_3$  is cyclopropyl,  $R_4$  is 1-methylimidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.12, 0.43, 0.68, 0.90–0.97, 1.36, 1.71, 2.60, 3.12, 3.67, 6.88, 7.06, 7.24, 7.51, 7.65.

HRMS found: 488.2225.

Anal. found: C, 61.25; H, 6.94; N, 8.42.

#### **EXAMPLE 49**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-8-quinolinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is quinolin-8-yl) Refer to Chart

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 8-quinolinesulfonyl chloride of formula D-7 wherein  $R_4$  is quinolin-8-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ-0.14, 0.01, 0.35, 0.89, 1.35, 1.63, 2.52, 2.94, 6.79, 6.94, 7.09, 7.64, 8.12, 8.28, 8.41, 9.13

HRMS found: 535.2256

Anal. found: C, 67.58; H, 6.53; N, 5.11.

# EXAMPLE 50

4-Cyano-N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is 4-cyanophenyl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 4-cyano-benzenesulfonyl chloride of formula D-7 wherein  $R_4$  is 4-cyanophenyl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.13, 0.44, 0.62, 0.91, 1.19, 1.67, 2.57, 3.14, 6.80, 7.12, 7.25, 7.83.

HRMS found: 509.2096

Anal. found: C, 65.86; H, 6.39; N, 5.48.

# **EXAMPLE 51**

4-Fluoro-N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is 4-fluorophenyl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 4-fluoro-benzenesulfonyl chloride of formula D-7 wherein  $R_4$  is 4-fluorophenyl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:  $^{1}$ H NMR (300 MHz, CD<sub>3</sub>OD)  $\delta$ 0.11, 0.43, 0.62, 0.92, 1.34, 1.65, 2.57, 3.13, 6.79, 7.03–7.24, 7.75.

HRMS found: 502.2063.

Anal. found: C, 63.96; H, 6.29; N, 2.71.

#### PREPARATION 21

Chiral HPLC resolution of 4-Hydroxy-3-[1-(3-nitrophenyl)-propyl]-5,6-dibydro-6,6-dipropyl-2H-pyran-2-one (Formula D-4:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl) Refer to Chart D

A solution of the title product of Preparation 18 (Formula D-4: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) (30 mg/mL) in 15% isopropyl alcohol in hexane is chromatographed on a 2.0×25 cm (R,R) Whelk-O 1 (Regis technologies, Inc., Morton Grove, Ill. 60053) column at 1 mL per injection 15 using an automated chromatographic system. The eluant is monitored at 270 nM and appropriate fractions from multiple injections combined and concentrated in vacuo to give tan oils. Fractions from multiple injections are analyzed on a 0.46×25 cm (S,S) Whelk-O 1 column with the same 20 solvent at 1.0 mL/min. The first peak from the 1.0 cm column is >99% ee (Rt is min) and the latter peak is 92% ee (Rt is min). Prior to further use, the resolved materials are subjected to flash chromatography on silica gel with 3:1 hexanes/EtOAc. The resolved materials are converted to the 25 amines of Formula D-5 using the conditions described in Preparation 20.

Physical characteristics are as follows:

The resolved materials were found to exhibit identical <sup>1</sup>H NMR and tlc behavior as the racemic material.

#### **EXAMPLE 52**

(R or S)-N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipropyl-6,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-8-quinolinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is quinolin-8-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 21 (Formula D-5: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 8-quinolinesulfonyl chloride of formula D-7 wherein R<sub>4</sub> is quinolin-8-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR and the behavior is identical to that of racemic

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.67, 0.85, 1.27, 1.54, 2.01, 3.73, 6.78, 6.90, 7.04, 7.57, 8.12, 8.29, 8.38, 9.13. MS m/e (rel%): 523 (100), 524 (34), 129 (11), 525 (11), 522 (10), 130 (7), 139 (5), 134 (4).

### **EXAMPLE 53**

(R or S)-N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 1-methylimidazol-4-yl) Refer to 55 Chart D

The title compound is prepared from the amine of Preparation 21 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the 60 general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR and tlc behavior is identical to racemic mixture. 65 <sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.88, 1.32, 1.64, 1.93, 2.16, 2.56, 3.68, 3.91, 6.87, 7.03, 7.14, 7.53, 7.64. 70

MS m/e (rel%): 476 (100), 477 (28), 139 (14), 492 (12), 134 (11), 278 (10), 478 (10), 83 (9), 552 (8), 145 (7).

#### **EXAMPLE 54**

(S or R)-N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-8-quinolinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is quinolin-8-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 21 (Formula D-5: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) and 8-quinolinesulfonyl chloride of formula D-7 wherein R<sub>4</sub> is quinolin-8-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR and tlc behavior is identical to that of racemic mixture.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.67, 0.85, 1.27, 1.54, 2.01, 3.73, 6.78, 6.90, 7.04, 7.57, 8.12, 8.29, 8.38, 9.13.

MS m/e (rel%): 523 (100), 524 (34), 522 (24), 539 (13), 525 (10), 129 (10), 130 (5), 134 (5), 128 (5), 540 (5).

#### **EXAMPLE 55**

(S or R)-N-[3-[1-(4-hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 1-methylimidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 21 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR and tle behavior is identical to racemic mixture. <sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.88, 1.32, 1.64, 1.93, 2.16, 2.56, 3.68, 3.91, 6.87, 7.03, 7.14, 7.53, 7.64.

MS m/e (rel%): 476 (100), 477 (28), 139 (19), 490 (15), 498 (14), 83 (12), 478 (9), 55 (9), 145 (9), 134 (7).

### **EXAMPLE 56**

(R or S)-N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is 1-methylimidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 21 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

<sup>1</sup>H NMR and tlc behavior is identical to racemic mixture. <sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.12, 0.43, 0.68, 0.90–0.97, 1.36, 1.71, 2.60, 3.12, 3.67, 6.88, 7.06, 7.24, 7.51, 7.65.

MS m/e (rel%): 488 (100), 489 (30), 139 (18), 145 (14), 490 (10), 55 (10), 83 (9), 564 (7), 146 (7), 510 (7).

#### **EXAMPLE 57**

(S or R)-N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipropyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is

propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is cyclopropyl, R<sub>4</sub> is 1-methylimidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine Preparation 21 (Formula D-5:  $R_1$  is propyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 4% MeOH/EtOAc.

Physical characteristics are as follows:

 $^{1}$ H NMR and tlc behavior is identical to racemic mixture.  $^{1}$ H NMR (300 MHz, CD<sub>3</sub>OD)  $\delta$ 0.12, 0.43, 0.68, 0.90–0.97, 1.36, 1.71, 2.60, 3.12, 3.67, 6.88, 7.06, 7.24, 7.51, 7.65.

MS m/e (rel%): 488 (100), 489 (29), 139 (18), 145 (16), 15 83 (10), 55 (10), 490 (10), 510 (8), 146 (8), 144 (7).

#### **EXAMPLE 58**

4-Cyano-N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is 4-cyanophenyl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is 25 cyclopropyl) and 4-fluoro-benzenesulfonyl chloride of formula D-7 wherein  $R_4$  is 4-cyanophenyl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.11, 0.42, 0.61, 0.95, 1.24, 1.74–2.00, 2.61–2.73, 3.30, 6.83–7.23, 7.71–7.84.

HRMS found: 571.2267

# **EXAMPLE 59**

4-Fluoro-N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is 4-fluorophenyl) Refer to 40 Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 4-fluoro-benzenesulfonyl chloride of formula D-7 wherein  $R_4$  is 4-fluorophenyl, using the general 45 procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.11, 0.43, 0.67, 0.96, 1.41, 1.67–2.13, 2.62, 3.16, 6.84, 7.02–7.31, 7.72.

HRMS found: 564.2211.

### **EXAMPLE 60**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-8-quinolinesulfonamide (Formula D-6:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is quinolin-8-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 8-quinolinesulfonyl chloride of formula D-7 wherein  $R_4$  is quinolin-8-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

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Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ-0.13, 0.01, 0.35, 0.93, 1.46, 1.54, 1.58-2.06, 2.56, 2.96, 6.81-7.23, 7.50-7.68, 8.08, 8.24, 8.37, 9.12.

HRMS found: 597.2398

#### **EXAMPLE 61**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl,  $R_4$  is 1-methyl-imidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is cyclopropyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazol-4-yl using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.13, 0.42, 0.67, 0.95, 1.44, 1.68–2.13, 2.56, 3.17, 6.91, 7.01–7.33, 7.52, 7.63.

HRMS found: 550.2370.

### **EXAMPLE 62**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-cyclopropylmethyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is phenethyl,  $R_2$  is phenethyl,  $R_3$  is cyclopropyl,  $R_4$  is 1-methyl-imidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is cyclopropyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein R<sub>4</sub> is 1-methylimidazol-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.13, 0.42, 0.68, 1.73, 1.88–2.17, 2.68, 3.19, 3.64, 6.93, 7.02–7.31, 7.52, 7.64.

HRMS found: 612.2530.

### **EXAMPLE 63**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipentyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6:  $R_1$  is pentyl,  $R_2$  is pentyl,  $R_3$  is ethyl,  $R_4$  is 1-methyl-imidazol-4-yl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is pentyl,  $R_2$  is pentyl,  $R_3$  is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 1-methylimidazole-4-yl, using the general procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) 80.87, 1.25, 1.55–1.68, 1.92, 2.13, 2.57, 3.66, 3.93, 6.86, 7.03, 7.16, 7.55, 7.63.

#### **EXAMPLE 64**

60 4-Cyano-N-[3-[1-(4-Hydroxy-2-oxo-6,6-dipentyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-benzenesulfonamide (Formula D-6: R<sub>1</sub> is pentyl, R<sub>2</sub> is pentyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 4-cyanophenyl) Refer to Chart D

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is pentyl,  $R_2$  is pentyl,  $R_3$  is ethyl) and 1-methyl-imidazole-4-sulfonyl chloride of formula D-7 wherein  $R_4$  is 4-cyanophenyl, using the general

procedure for sulfonylation of Example 40 to yield the title compound as a white amorphous solid after flash chromatography with 5% EtOAc/CH2Cl2.

Physical characteristics are as follows:

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) δ0.86, 1.23, 1.52–1.67, 5 1.93, 2.14, 2.56, 3.93, 6.80, 7.05, 7.18, 7.80, 7.86.

### EXAMPLES 65-93

Using the general procedure of Example 40, but substituting the appropriate reactants, the following compounds of the present invention are prepared:

#### **EXAMPLE 65**

N-[3-[1(R or S)-(6(R or S)-4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1- 15 methyl-1H-imidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R2 is propyl, R3 is ethyl, R4 is 1-methylimidazol-4-yl) Refer to Chart D

#### **EXAMPLE 66**

N-[3-[1(R or S)-(6(S or R)-4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1methyl-1H-imidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R2 is propyl, R3 is ethyl, R4 is 1-methylimidazol-4-yl) Refer to Chart D

### **EXAMPLE 67**

N-[3-[1(S or R)-(6(R or S)-4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1methyl-1H-imidazole-4-sulfonamide (Formula D-6: R, is 30 phenethyl, R2 is propyl, R3 is ethyl, R4 is 1-methylimidazol-4-yl) Refer to Chart D

### **EXAMPLE 68**

N-[3-[1(S or R)-(6(S or R)-4-Hydroxy-2-oxo-6-phenethyl- 35 R<sub>4</sub> is 2-pyrimidyl) Refer to Chart D 6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1methyl-1H-imidazole-4-sulfonamide (Formula D-6: R, is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 1-methylimidazol-4-yl) Refer to Chart D

### **EXAMPLE 69**

N-[3-[t-Butyl-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6dihydro-2H-pyran-3-yl)-methyl]-phenyl]-1-methyl-1Himidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is t-butyl, R<sub>4</sub> is 1-methylimidazol-4-yl) Refer 45 to Chart D

### **EXAMPLE 70**

4-Cyano-N-[3-[t-butyl-(4-hydroxy-2-oxo-6-phenethyl-6propyl-5,6-dihydro-2H-pyran-3-yl)-methyl]-phenyl]benzenesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is t-butyl, R<sub>4</sub> is 4-cyanophenyl) Refer to Chart D

#### **EXAMPLE 71**

4-Fluoro-N-[3-[t-butyl-(4-hydroxy-2-oxo-6-phenethyl-6propyl-5,6-dihydro-2H-pyran-3-yl)-methyl]-phenyl]- 55 benzenesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is t-butyl, R<sub>4</sub> is 4-fluorophenyl) Refer to Chart D

### **EXAMPLE 72**

N-[3-[t-Butyl-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-60 dihydro-2H-pyran-3-yl)-methyl]-phenyl]-8quinolinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is t-butyl, R<sub>4</sub> is quinolin-8-yl) Refer to Chart D

#### **EXAMPLE 73**

N-[3-[1-(6-(2-(1-Methyl-1H-imidazole-4-sulfonylamino)ethyl)-4-hydroxy-2-oxo-6-propyl-5,6-dihydro-2H-pyran-3-

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yl)-propyl]-phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is 2-(1-methylimidazole-4sulfonylamino)-ethyl, R2 is propyl, R3 is ethyl, R4 is 1-methylimidazol-4-yl) Refer to Chart D

#### **EXAMPLE 74**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-pyridinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 2-pyridyl) Refer to Chart D

#### EXAMPLE 75

4-Cyano-N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2pyridinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 4-cyano-2-pyridyl) Refer to Chart D

#### **EXAMPLE 76**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-quinolinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is quinolin-2-yl) Refer to Chart D

#### **EXAMPLE 77**

25 2-Hydroxy-N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6dihydro-2H-pyran-3-yl)-propyl]-phenyl]benzenesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 2-hydroxyphenyl) Refer to Chart D

#### **EXAMPLE 78**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-pyrimidinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl,

# **EXAMPLE 79**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-quinazolinesulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is quinazolin-2-yl) Refer to Chart D

### **EXAMPLE 80**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-7H-purine-6-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 7H-purin-6-yl) Refer to Chart D

#### **EXAMPLE 81**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1H-imidazole-2sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 1H-imidazol-2-yl) Refer to Chart D

### **EXAMPLE 82**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1H-benzimidazole-2sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 1H-benzimidazol-2-yl) Refer to Chart D

# **EXAMPLE 83**

N-[3-[1-(4-Hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-thiazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is thiazol-2-yl) Refer to Chart D

#### **EXAMPLE 84**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-

pyridinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 2-pyridyl) Refer to Chart D

### **EXAMPLE 85**

4-Cyano-N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-pyridinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 4-cyano-2-pyridyl) Refer to Chart D

#### **EXAMPLE 86**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-quinolinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is quinolin-2-yl) Refer to Chart D 15

# **EXAMPLE 87**

2-Hydroxy-N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 2-hydroxyphenyl) Refer to Chart D

### **EXAMPLE 88**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-pyrimidinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 2-pyrimidyl) Refer to Chart D

### **EXAMPLE 89**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-2-quinazolinesulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is quinazolin-2-yl) Refer to Chart D

# **EXAMPLE 90**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-7H-purine-6-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 7H-purin-6-yl) Refer to Chart D

### **EXAMPLE 91**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1H-imidazole-2-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 1H-imidazol-2-yl) Refer to Chart D

### **EXAMPLE 92**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-1H-  $^{50}$  benzimidazole-2-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is 1H-benzimidazol-2-yl) Refer to Chart D

### **EXAMPLE 93**

N-[3-[1-(4-Hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-thiazole-4-sulfonamide (Formula D-6:  $R_1$  is propyl,  $R_2$  is phenethyl,  $R_3$  is ethyl,  $R_4$  is thiazol-2-yl) Refer to Chart D

### **EXAMPLE 93A**

4-Fluoro-N-[3-[1-(4-hydroxy-2-oxo-6-phenethyl-6-propyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-benzenesulfonamide (Formula D-6:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 4-fluorophenyl) Refer to Chart D 65

The title compound is prepared from the amine of Preparation 20 (Formula D-5:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is

ethyl) and 4-fluoro benzenesulfonyl chloride using the general procedure for sulfonylation of Example 40 to yield the title compound as an off-white amorphous solid after flash chromatography with 5% EtOAc/CH<sub>2</sub>Cl<sub>2</sub>.

Physical characteristics are as follows:

<sup>1</sup>H NMR complicated by presence of diastereomers.

<sup>1</sup>H NMR (300 MHz, CD<sub>3</sub>OD) 80.75–0.96, 1.31–1.48, 1.57–2.01, 2.09–2.22, 2.48–2.71, 3.92, 3.94, 6.86–7.24, 7.72.

#### PREPARATION 22

6-(2-Cyclopropyl-1-cyclopropylmethyl-ethyl)-4-hydroxy-2H-pyran-2-one (Formula E-2) Refer to Chart E

To a cold (-78° C.) stirred solution of 1.5 ml of diisopropylamine in 9 ml of dry tetrahydrofuran, under argon, is added 6.2 ml of a 1.6M solution of n-butyllithium in hexane. The solution is warmed to 0° C. and then treated with a solution of 378 mg of commercially available 4-hydroxy-6-methyl-2-pyrone of formula E-1 in 8 ml of hexamethylphosphoramide. After 30 minutes at 0° C., 0.32 ml of bromomethylcyclopropane is added; after another ten minutes, a second portion of the same amount is added. The reaction is stirred, allowed to warm to room temperature overnight, and is then partitioned between ethyl acetate and excess dilute hydrochloric acid. The organic phase is washed with brine, dried over magnesium sulfate, and concentrated under reduced pressure. The residue is flash chromatographed on silica gel 60 (230-400 mesh) using 25% ethyl acetate in dichloromethane containing 1% acetic acid to provide 371 mg of the title compound, along with 206 mg of monoalkylated material.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.0, 0.4, 0.6, 1.5, 1.6, 2.2, 5.6, 6.1, 7.2–7.3, 11.5:

EI MS m/z=234;

TLC  $R_f$  0.29 (25% ethyl acetate in dichloromethane containing 1% acetic acid).

### PREPARATION 23

3-(\alpha-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-6-(2-cyclopropyl-1-cyclopropylmethyl-ethyl)-4-hydroxy-2H-pyran-2-one (Formula E-3) Refer to Chart E

A mixture of 367 mg of the title compound of Preparation 22, 470 mg of the title compound of Preparation F-5, 60 mg of p-toluenesulfonic acid monohydrate, and 1 g of 3 Å molecular sieves in 5 ml of benzene is heated with stirring overnight under argon. The mixture is diluted with dichloromethane and ether and filtered through a pad of sodium sulfate. The solvent is removed under reduced pressure and the residue is flash chromatographed on silica gel 60 (230–400 mesh) using 5–20% ethyl acetate in dichloromethane to afford 399 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ-0.06, 0.3, 0.5, 1.4, 1.5, 2.5, 3.5, 5.1, 7.2-7.4; EI HRMS m/z=513.2513;

TLC R<sub>f</sub> 0.28 (5% ethyl acetate in dichloromethane).

#### PREPARATION 24

3-(\alpha-Cyclopropyl-meta-aminobenzyl)-6-(2-cyclopropyl-1-cyclopropylmethyl-ethyl)-4-hydroxy-2H-pyran-2-one (Formula E-4) Refer to Chart E

A mixture of 391 mg of the title compound of Preparation 23 and 100 mg of 5% palladium on carbon in 10 ml of methanol is shaken overnight under 40 psi of hydrogen. The mixture is then filtered through Celite, and the filtrate is concentrated under reduced pressure to provide 280 mg of the title compound.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.0, 0.2–0.7, 1.4, 1.6, 1.8, 2.6, 6.8, 7.2–7.4; TLC  $R_f$  0.38 (30% ethyl acetate in dichloromethane.)

#### **EXAMPLE 94**

N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] -methyl}-phenyl)-4-fluoro-benzenesulfonamide (Formula E-6) Refer to Chart E

ration 24 and 24  $\mu$ L of pyridine in 0.5 mL of dichloromethane is added 29 mg of 4-fluorobenzenesulfonyl chloride. After stirring overnight, the solution is diluted with ethyl acetate and washed with dilute aqueous hydrochloric acid, brine, dried over sodium sulfate, and concentrated 15 under reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) using 10% ethyl acetate in dichloromethane to give 56 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ-0.07, 0.13, 0.33, 0.54, 1.39, 1.51, 1.72, 2.55, 3.39, 6.12, 6.87, 7.00, 7.08, 7.19, 7.27, 7.72, 9.72;

EI-MS: [M+]=537.1977 found.

### **EXAMPLES 95-97**

Following the procedure described above and using starting materials and reagents known and available to one of ordinary skill in organic synthesis, the following additional compounds are prepared:

#### **EXAMPLE 95**

4-Cyano-N-(3-{cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] -methyl}-phenyl)-benzenesulfonamide (Formula E-7) Refer 35 to Chart E

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ-0.03, 0.13, 0.23, 0.36, 0.44, 0.57, 1.41, 1.58, 1.75, 2.57, 3.32, 5.98, 6.89, 7.11, 7.21, 7.68, 7.82; EI-MS: [M+]=544.2035 found.

### **EXAMPLE 96**

N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] Refer to Chart E

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ-0.07, 0.18, 0.37, 0.54, 1.37, 1.51, 2.53, 3.31, 5.96, 6.87, 7.00, 7.13, 7.48, 7.54, 7.92, 8.23, 9.07;

EI-MS: [M+]=570.2188 found.

#### EXAMPLE 97

N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] 55 -methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula E-9) Refer to Chart E

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>---CD<sub>3</sub>OD) δ-0.08, 0.13, 0.33, 0.56, 7.5;

EI-MS: [M+]=523.2142 found.

### **EXAMPLE 98**

Chiral HPLC Separation of N-(3-{Cyclopropyl-[6-(2-65 cyclopropyl-1-cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole78

4-sulfonamide (Formula E-9) to give (R or S)-N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1-cyclopropylmethylethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula E-10) and (R or S)-N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] -methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula E-11) Refer to Chart E

A stock sample of the title compound of Example 97 (3 To a mixture of 57 mg of the title compound of Prepa- 10 mg/ml) in 5.0 mL each of mobile phase (30% isopropanol, 0.1% acetic acid, and 0.2% water in hexane) and isopropanol is prepared. The stock sample is filtered through a 0.45 micron syringe filter and washed with ethanol to give 14.0 mL of clear filtrate. This solution is chromatographed on a 2.0×2.5 cm (R,R) Whelk-O 1 (Regis Technologies, Inc., Morton Grove, Ill. 60053) column at 3.50 mL per injection using an automated chromatographic system. The eluant is monitored and the pools corresponding to the desired peaks from multiple injections are combined, concentrated under reduced pressure and azeotroped with toluene. The residues are dissolved in methanol, filtered through a syringe filter and the filtrates concentrated under reduced pressure to give the title compounds (>95% pure):

(R or S)-N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1-25 cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl -methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula E-10)

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ-0.07, 0.14, 0.34, 0.57, 30 1.32, 1.55, 1.75, 2.51, 3.24, 3.60, 5.87, 6.85, 7.03, 7.15, 7.27, 7.37;

EI-MS: [M+]=523.2149 found.

(R or S)-N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] -methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula E-11)

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD)  $\delta$ -0.07, 0.14, 0.33, 0.55, 1.33, 1.56, 1.75, 2.51, 3.23, 3.60, 5.88, 6.86, 7.03, 7.14, 7.27, 40 7.38;

EI-MS: [M+]=523.2137 found.

#### EXAMPLES 99-103

Following the procedure described above and using start--methyl}-phenyl)-8-quinolinesulfonamide (Formula E-8) 45 ing materials and reagents known and available to one of ordinary skill in organic synthesis, the following additional compounds are prepared:

#### **EXAMPLE 99**

N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] -methyl}-phenyl)-2-pyridinesulfonamide (Formula E-12)

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD)  $\delta$ -0.05, 0.07, 0.17, 0.34, 0.55, 1.35, 1.55, 1.7, 2.5, 3.24, 5.86, 6.90, 7.03, 7.15, 7.39, 7.78, 8.60;

EI-MS: [M+]=520;

TLC R<sub>c</sub> 0.35 (25% ethyl acetate in dichloromethane).

### **EXAMPLE 100**

1.37, 1.51, 1.73, 2.54, 3.21, 3.60, 5.95, 6.82, 7.0, 7.19, 7.37, 60 N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl] -methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide (Formula E-13)

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ-0.05, 0.15, 0.35, 0.56, 1.35, 1.55, 1.75, 2.53, 3.23, 3.39, 5.89, 6.81, 6.90, 6.97, 7.09,

EI-MS: [M+]=523; TLC R<sub>f</sub> 0.31 (5% methanol in dichloromethane).

### **EXAMPLE 101**

N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1-5 cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-benzo-imidazole-2-sulfonamide (Formula E-14)

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ-0.07, 0.1, 0.15, 0.35, <sub>10</sub> 0.56, 1.38, 1.58, 1.65, 2.55, 3.28, 5.95, 6.73, 6.96, 7.10, 7.28, 7.58;

FAB-MS: [M+H]=560.2220 found.

### **EXAMPLE 102**

N-(3-{Cyclopropyl-[6-(2-cyclopropyl-1-cyclopropyl-1-cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide (Formula E-15)

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) 80.0, 0.25, 0.4, 0.6, 1.4, 1.6, 1.65, 2.6, 3.35, 6.0, 6.8, 7.0, 7.2, 7.4;

EI-MS: [M+]=509;

TLC R<sub>f</sub> 0.25 (5% methanol in dichloromethane).

### **EXAMPLE 103**

 $N-(3-\{C\ y\ c\ l\ o\ p\ r\ o\ p\ y\ l-[6-(2-c\ y\ c\ l\ o\ p\ r\ o\ p\ y\ l-1-cyclopropylmethyl-ethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl\}-phenyl)-2-quinolinesulfonamide (Formula E-16)$ 

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.0, 0.2, 0.4, 0.6, 1.4, 1.6, 1.7, 2.6, <sup>30</sup> 3.3, 6.0, 7.0–7.2, 7.3, 7.7, 7.8–8.0, 8.2, 8.3;

EI-MS: [M+]=570;

TLC R<sub>f</sub> 0.53 (5% methanol in dichloromethane).

### PREPARATION 25

Cyclopropyl meta-nitrophenyl ketone (Formula F-2) Refer to Chart F

A 250 ml three necked flask fitted with thermometer and addition funnel is charged with 130 ml of fuming 90% nitric acid and cooled to -10° C. Into the stirred liquid is added 40 dropwise 21 ml of commercially available cyclopropyl phenyl ketone of formula F-1. The rate of addition is regulated to maintain the reaction temperature at about -10° C. Upon completion of addition, the resulting clear yellow solution is stirred for another 10 minutes at -10° C., then 45 poured into 1 L of crushed ice. The precipitated solid is extracted with 700 ml of toluene, and the extract is washed twice with 5% sodium hydroxide solution, once with brine, and dried over magnesium sulfate. The solvent is removed under reduced pressure and the residue is recrystallized from 50 methanol at -25° C. to give 14.6 g of the title compound as dense, pale yellow prisms. The mother liquor contained substantial amounts of the ortho isomer.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ1.2, 1.3, 2.7, 7.70, 8.3, 8.4, 8.85;

IR 1664, 1529, 1352, 1225, 1082, 1017, 852, 689 cm<sup>-1</sup>; Anal. Found: C, 62.89; H, 4.73; N, 7.32;

EI MS m/z 191;

TLC R, 0.32 (25% ethyl acetate in hexane).

# PREPARATION 26

meta-Aminophenyl cyclopropyl ketone (Formula F-3) Refer to Chart F

A solution of 5.76 g of the title compound of Preparation 25 is prepared with the aid of heat in 100 ml of methanol. 65 To this is added 450 mg of 5% platinum on carbon catalyst, and the mixture is stirred vigorously under 1 atmosphere of

80

hydrogen. After 5 hours, the mixture is filtered through a pad of Celite and the filtrate concentrated under reduced pressure to afford 4.89 g of the title compound as a greenish oil.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 1.0, 1.2, 2.6, 3.9, 6.8, 7.2, 7.4; TLC  $R_f$  0.50 (80% ethyl acetate in hexane).

### PREPARATION 27

meta-Benzyloxycarbonylaminophenyl cyclopropyl ketone (Formula F-4) Refer to Chart F

To a cold (0° C.), stirred solution of 4.89 g of the title compound of Preparation 26 and 6.3 ml of diisopropylethylamine in 90 ml of dichloromethane is added dropwise 4.7 ml of benzyl chloroformate. The completed solution is allowed to warm to room temperature. After 4 hours, the mixture is washed with dilute hydrochloric acid, and the aqueous phase extracted with two additional portions of dichloromethane. The combined organic phase is dried over magnesium sulfate and concentrated under reduced pressure to a yellow solid. This is triturated with two 30 ml portions of hexane, these being discarded, and the remaining solid is dried under vacuum to afford 8.74 g of the title compound. Physical characteristics are as follows:

TLC Rf 0.45 (5% ethyl acetate in dichloromethane).

#### PREPARATION 28

meta-Benzyloxycarbonylaminophenyl cyclopropyl carbinol (Formula F-5) Refer to Chart F

To a stirred solution of 8.74 g of compound F-4 of Preparation 27 in 100 ml of tetrahydrofuran and 100 ml of ethanol is added, in portions, 4.5 g of sodium borohydride. After 3 hours at room temperature, the mixture is cooled in ice for the addition of 100 ml of 1N hydrochloric acid. The mixture is thrice extracted with dichloromethane, and the combined extract dried over magnesium sulfate. Solvent is removed under reduced pressure and the residue flash chromatographed on silica gel 60 (230–400 mesh) using 40% ethyl acetate in hexane to provide 8.48 g of the title compound as a white crystalline solid. This is optionally recrystallized from ethyl acetate-hexane.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.3–0.6, 1.1, 2.35, 3.92, 5.17, 7.1, 7.2–7.4; IR 1693, 1599, 1559, 1449, 1235, 1054, 697 cm<sup>-1</sup>; Anal. Found: C, 72.57; H, 6.51; N, 4.61;

## PREPARATION 29

4-Hydroxy-6-[3-(2-methoxy-ethoxy)-propyl]-pyran-2-one (Formula G-1) Refer to Chart G

To a flame-dried flask under an argon atmosphere is added 2.80 mL of disopropylamine and 20.0 mL of dry tetrahydrofuran. The solution is cooled to -78° C. and treated with 12.5 mL (1.6M in hexane) of n-butyllithium. The solution is warmed to 0° C. for 30 minutes, then treated with 5.0 mL of dry hexamethylphosphoramide. The lithium diisopropylamide solution is then treated with 1.20 g of commercially available 4-hydroxy-6-methyl-2-pyrone of formula G-0 as a solution in 16 mL of dry tetrahydrofuran and 14 mL of dry hexamethylphosphoramide. After 30 minutes the mixture is treated with 2.30 g of 2-(2-methoxy-ethoxy)-ethyl iodide as a solution in 12 mL of dry tetrahydrofuran. The mixture is stirred 1 hour at 0° C. and then warmed to room temperature. After 1 hour the reaction is quenched with excess 1N aqueous hydrochloric acid. The mixture is concentrated under reduced pressure and partioned between dichloromethane and water. The aqueous phase is extracted with sufficient volumes of dichloromethane to remove the title compound. The combined organic extracts are washed with brine, dried over magnesium sulfate and concentrated under

reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with 30% ethyl acetate in dichloromethane containing 3% acetic acid to 80% ethyl acetate in dichloromethane containing 5% acetic acid to give 1.34 g of the title compound. 5

Physical characteristics are as follows:

 $^{1}$ H-NMR (CDCl<sub>3</sub>) δ1.93, 2.54, 3.39, 3.55, 5.55, 5.90; TLC R<sub>f</sub> 0.26 (50% ethyl acetate in dichloromethane containing 5% acetic acid).

#### PREPARATION 30

3-(\alpha-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4-hydroxy-6-[3-(2-methoxy-ethoxy)-propyl]-pyran-2-one (Formula G-2) Refer to Chart G

A mixture of 146 mg of the title compound of Preparation 29, 340 mg of the title compound of Preparation 28, prepared as described in Chart F, 25 mg of p-toluenesulfonic acid monohydrate, and 0.5 g of 3 Å molecular sieves in 5 mL of dichloromethane is heated overnight with stirring. The mixture is cooled and the solvent removed under reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 5% to 10% methanol in ethyl acetate to give 129 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.25, 0.45, 0.67, 1.77, 2.39, 3.38, 3.51, 5.13, 5.84, 7.17, 7.32, 7.42;

EI-MS: [M+]=507.2257 found;

TLC R<sub>f</sub> 0.28 (50% ethyl acetate in dichloromethane).

### PREPARATION 31

 $3-(\alpha-Cyclopropyl-meta-aminobenzyl)-4-hydroxy-6-[3-(2-methoxy-ethoxy)-propyl]-pyran-2-one (Formula G-3) Refer to Chart G$ 

A mixture of 124 mg of the title compound of Preparation 30 and 35 mg of 5% palladium on charcoal in 5 mL of ethanol is shaken overnight under 50 psi of hydrogen. The mixture is filtered through Celite with ethanol washes of the filter cake. The filtrates are combined and the solvent removed under reduced pressure to give 92 mg of the title compound. Physical characteristics are as follows:

TLC  $R_f$  0.12 (ethyl acetate).

### **EXAMPLE 104**

N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{2-methoxy-ethoxy}-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula G-4) Refer to Chart G

To a mixture of 37 mg of the title compound of Preparation 31 and 18  $\mu$ L of pyridine in 0.5 mL of dichloromethane is added 20 mg of 1-methylimidazole-4-sulfonyl chloride. After stirring overnight, the solvent is removed under reduced pressure. The residue is azeotroped with toluene and is then purified by flash column chromatography on silica gel 60 (230–400 mesh) using 2% to 8% methanol in dichloromethane to give 32 mg of the title compound. 55

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) 80.1, 0.24, 0.45, 0.65, 1.75, 1.85, 2.46, 3.30, 3.34, 3.50, 5.98, 6.98, 7.08, 7.19, 7.29, 7.42;

EI-MS: [M+]=517.1874 found;

TLC R<sub>f</sub> 0.22 (5% methanol in dichloromethane).

### PREPARATION 32

3-(\alpha-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4-hydroxy-6-methyl-pyran-2-one (Formula H-1) Refer to Chart H

A mixture of 493 mg of commercially available 4-hydroxy-6-methyl-2-pyrone of formula H-0, 592 mg of 82

the title compound of Preparation 28, prepared as described in Chart F and 56 mg of p-toluenesulfonic acid monohydrate in 20 mL of dichloromethane is heated to reflux through an addition funnel containing 3 Å molecular sieves for 6 hours. The mixture is cooled and the solvent removed under reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 60% to 100% ethyl acetate in dichloromethane to give 470 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) 80.23, 0.43, 0.66, 1.78, 3.41, 5.09, 5.89, 7.00, 7.14, 7.29, 7.37, 10.1;

EI-MS: [M+]=405;

TLC R<sub>f</sub> 0.52 (ethyl acetate).

#### PREPARATION 33

3-(α-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4-hydroxy-6-propyl-pyran-2-one (Formula H-2) Refer to Chart H

To a flame-dried flask under an argon atmosphere is added 0.45 mL of diisopropylamine and 3.0 mL of dry tetrahydrofuran. The solution is cooled to -78° C. and treated with 2.0 mL (1.6M in hexane) of n-butyllithium. The solution is warmed to 0° C. for 15 minutes, then cooled to -78° C. The lithium diisopropylamide solution is treated with 405 mg of the title compound of Preparation 32 as a solution in 4 mL of dry tetrahydrofuran. After 1 hour at -78° C. the mixture is treated with 85  $\mu$ L of ethyl bromide. The mixture is then stirred at -78° C. for 3 hours. The reaction is quenched with excess 1N aqueous hydrochloric acid. The mixture is warmed and partioned between ethyl acetate and phosphate buffer. The aqueous phase is extracted twice with ethyl acetate. The combined organic extracts are dried over magnesium sulfate and concentrated under reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with 10% to 20% ethyl acetate in dichloromethane to give 277 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.24, 0.45, 0.65, 0.88, 1.55, 1.79, 2.28, 3.42, 5.10, 5.95, 6.89, 7.15, 7.3, 10.0;

EI-MS: [M+]=433;

TLC  $R_f$  0.33 (10% ethyl acetate in dichloromethane).

#### PREPARATION 34

 45 3-(α-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-6-[1-ethyl-3-(2-methoxy-ethoxy)-propyl]-4-hydroxy-pyran-2-one (Formula H-3) Refer to Chart H

To a flame-dried flask under an argon atmosphere is added 0.30 mL of diisopropylamine and 2.0 mL of dry tetrahydrofuran. The solution is cooled to -78° C. and treated with 1.3 mL (1.6M in hexane) of n-butyllithium. The solution is warmed to 0° C. for 15 minutes, then cooled to ~78° C. The lithium diisopropylamide solution is treated with 277 mg of the title compound of Preparation 33 as a solution in 3 mL of dry tetrahydrofuran. After 1 hour at -78° C., the mixture is treated with 180 mg of 2-(2-methoxy-ethoxy)-ethyl iodide in 3 mL of tetrahydrofuran. The mixture is then stirred at -78° C. for 3 hours. The reaction is quenched with excess 1N aqueous hydrochloric acid. The mixture is warmed and partioned between ethyl acetate and phosphate buffer. The aqueous phase is extracted thrice with ethyl acetate. The combined organic extracts are washed with brine, dried over magnesium sulfate and concentrated under reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with 25% to 40% ethyl acetate in dichloromethane to give 198 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.25, 0.46, 0.66, 0.75, 1.5, 1.76, 2.36, 3.4, 5.15, 5.96, 7.2, 7.3, 7.42, 10.0;

EI-MS: [M+]=535;

TLC R<sub>c</sub> 0.29 (25% ethyl acetate in dichloromethane).

#### PREPARATION 35

3-(α-Cyclopropyl-meta-aminobenzyl)-6-[1-ethyl-3-(2methoxy-ethoxy)-propyl]-4-hydroxy-pyran-2-one (Formula H-4) Refer to Chart H

A mixture of 180 mg of the title compound of Preparation 34 and 50 mg of 5% palladium on charcoal in 2 mL of ethanol is shaken overnight under 50 psi of hydrogen. The mixture is filtered through Celite with ethanol washes of the filter cake. The filtrates are combined and the solvent 15 removed under reduced pressure to give 127 mg of the title

Physical characteristics are as follows: TLC  $R_f$  0.19 (ethyl acetate).

### **EXAMPLE 105**

N-(3-{Cyclopropyl-[6-(1-ethyl-3-{2-methoxy-ethoxy}propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula H-5) Refer to Chart H

To a mixture of 32 mg of the title compound of Preparation 35 and 13  $\mu$ L of pyridine in 0.8 mL of dichloromethane is added 14.5 mg of 1-methylimidazole-4sulfonyl chloride. After stirring overnight, the solvent is removed under reduced pressure. The residue is azeotroped with toluene and is then purified by flash column chromatography on silica gel 60 (230-400 mesh) using 1% to 4% methanol in dichloromethane to give 38 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) 80.05, 0.25, 0.45, 0.65, 0.83, 1.5–2.0, <sup>35</sup> 2.45, 3.3–3.5, 3.62, 6.00, 6.99, 7.1–7.3, 7.48;

EI-MS: [M+]=545.2186 found;

TLC R<sub>f</sub> 0.24 (5% methanol in dichloromethane).

#### **EXAMPLE 106**

4-Cyano-N-(3-{cyclopropyl-[6-(1-ethyl-3-{2-methoxyethoxy}-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]methyl}-phenyl)-benzenesulfonamide

Following the procedure described above and using starting materials and reagents known and available to one of 45 ordinary skill in organic synthesis, the title compound is prepared.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.15, 0.25, 0.45, 0.65, 0.78, 1.2–1.8, 2.4, 3.3-3.6, 3.54, 5.89, 6.95, 7.1-7.3, 7.6-7.9;

FAB-MS: [M+H]=567.2176 found;

TLC R<sub>f</sub> 0.40 (50% ethyl acetate in dichloromethane).

# PREPARATION 36

3-(\alpha-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)- 55 6-(1-ethyl-3-hydroxy-propyl)-4-hydroxy-pyran-2-one (Formula I-1) Refer to Chart I

To a flame-dried flask under an argon atmosphere is added 0.46 mL of disopropylamine and 3.5 mL of dry tetrahydrofuran. The solution is cooled to -78° C. and treated with 2.0 60 mL (1.6M in hexane) of n-butyllithium. The solution is warmed to 0° C. for 20 minutes, then cooled to -78° C. The lithium diisopropylamide solution is treated with 433 mg of the title compound of Preparation 33 as a solution in 4 mL of dry tetrahydrofuran. After 1 hour at -78° C. the mixture 65 is treated with gaseous ethylene oxide for 5 minutes. The mixture is then stirred at -78° C. for 15 minutes. The

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reaction is quenched with excess 1N aqueous hydrochloric acid. The mixture is warmed and partioned between dichloromethane and phosphate buffer. The aqueous phase is extracted twice with dichloromethane. The combined organic extracts are washed with brine, dried over magnesium sulfate and concentrated under reduced pressure. The residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with 2% to 8% methanol in dichloromethane to give 144 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.22, 0.45, 0.65, 0.7, 1.3–1.7, 1.8, 2.25, 3.4, 5.1, 5.91, 7.1-7.4;

FAB-MS: [M+H]=478;

TLC R<sub>f</sub> 0.29 (5% methanol in dichloromethane).

#### PREPARATION 37

6-(3-Bromo-1-ethyl-propyl)-3-(α-cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4-hydroxy-pyran-2-one (Formula I-2) Refer to Chart I

To a stirring solution of 114 mg of the title compound of Preparation 36 in 3 mL of tetrahydrofuran is added 160 mg of triphenylphosphine and 200 mg of carbon tetrabromide. After 2 hours, the solvent is removed under reduced pressure and the residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with 70% to 100% diethyl ether in hexane to give 113 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.25, 0.35, 0.55, 0.65, 0.84, 1.5–2.2, 3.2, 3.35, 3.52, 5.16, 5.95, 6.79, 7.1–7.4;

FAB-MS: [M+H]=504.1404 found;

TLC  $R_f$  0.29 (75% diethyl ether in hexane).

#### PREPARATION 38

6-(3-Azido-1-ethyl-propyl)-3-(α-cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4-hydroxy-pyran-2-one (Formula I-3) Refer to Chart I

To a stirring solution of 113 mg of the title compound of Preparation 37 in 2.0 mL of ethanol is added 55 mg of sodium azide and 0.5 mL of water. The reaction mixture is heated overnight and then cooled. The solvent is removed under reduced pressure and the residue is purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with diethyl ether to give 89 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.23, 0.33, 0.51, 0.68, 0.82, 1.4–2.0, 2.33, 3.1-3.3, 3.5, 5.15, 5.94, 6.84, 7.1-7.4;

EI-MS: [M+]=502;

TLC R<sub>f</sub> 0.52 (10% ethyl acetate in dichloromethane).

#### PREPARATION 39

6-(3-Amino-1-ethyl-propyl)-3-(α-cyclopropyl-metaaminobenzyl)- 4-hydroxy-pyran-2-one (Formula I-4) Refer to Chart I

A mixture of 87 mg of the title compound of Preparation 38 and 35 mg of 5% palladium on charcoal in 4 mL of ethanol is shaken for 4 hours under 40 psi of hydrogen. The mixture is filtered through Celite with ethanol washes of the filter cake. The filtrates are combined and the solvent removed under reduced pressure to give 70 mg of the title compound as a mixture with 6-(3-amino-1-ethyl-propyl)-3-(α-cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4hydroxy-pyran-2-one.

Physical characteristics are as follows:

TLC  $R_f$  0.05 (5% methanol in dichloromethane).

### PREPARATION 40

3-(Cyclopropyl-{3-[1-methyl-1H-imidazole-4sulfonylamino]-phenyl}-methyl)-6-({1-ethyl-3-[1-methyl-

1H-imidazole-4-sulfonylamino]}-propyl)-2-oxo-2H-pyran-4-yl 1-methyl-1H-imidazole-4-sulfonate (Formula I-5) Refer to Chart I

To a mixture of 70 mg of the title compound of Preparation 39 and 6-(3-amino-1-ethyl-propyl)-3-( $\alpha$ -cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-4-hydroxy-pyran-2-one, also from Preparation 39, in 1.5 mL of dichloromethane is added 120  $\mu$ L of diisopropylethylamine and 92 mg of 1-methylimidazole-4-sulfonyl chloride. After stirring overnight, the solvent is removed under reduced pressure. The residue is azeotroped with toluene and is then purified by flash column chromatography on silica gel 60 (230-400 mesh) using 2% to 6% methanol in dichloromethane to give 49 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) 80.2–0.5, 0.75, 0.90, 1.4–2.0, 2.55, <sub>15</sub> 3.0–3.4, 3.6–3.7, 6.63, 7.0–7.7;

TLC R<sub>f</sub> 0.14 (5% methanol in dichloromethane).

#### **EXAMPLE 107**

N-(3-{Cyclopropyl-[6-(1-ethyl-3-{1-methyl-1H-imidazole-4-sulfonylamino}-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula I-6) Refer to Chart I

A solution of 49 mg of the title compound of Preparation 40 in 4 mL methanol containing ammonia is cooled to 0° C. and treated with gaseous ammonia. After 5 minutes ammonia introduction is ceased, the flask is tightly capped and warmed to room temperature. After standing overnight the solvent is removed under reduced pressure and the residue is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 3% to 9% methanol in dichloromethane to give 32 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) 80.2–0.5, 0.75, 0.90, 1.4–2.0, 2.55, 3.0–3.4, 3.6–3.7, 6.63, 7.0–7.7;

EI-MS: [M+]=523;

TLC  $R_f$  0.33 (5% methanol in dichloromethane).

### PREPARATION 41

3-(\alpha-Cyclopropyl-meta-aminobenzyl)-6-(1-ethyl-3-hydroxypropyl)-4-hydroxy-pyran-2-one (Formula J-1) Refer to Chart J

A mixture of 477 mg of the title compound of Preparation 36 and 150 mg of 5% palladium on charcoal in 10 mL of ethanol is shaken overnight under 45 psi of hydrogen. The mixture is filtered through Celite with ethanol washes of the filter cake. The filtrates are combined and the solvent is removed under reduced pressure to give 340 mg of the title compound.

Physical characteristics are as follows:

TLC R, 0.10 (5% methanol in dichloromethane).

### PREPARATION 42

6-(3-Bromo-1-ethyl-propyl)-3-(α-cyclopropyl-meta-aminobenzyl)-4-hydroxy-pyran-2-one (Formula J-2) Refer to Chart J

To a stirring solution of 340 mg of the title compound of Preparation 41 in 7 mL of tetrahydrofuran is added 525 mg of triphenylphosphine and 663 mg of carbon tetrabromide. After 30 minutes, the solvent is removed under reduced pressure and the residue is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 30% to 50% ethyl acetate in dichloromethane to give 228 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>) δ0.19, 0.42, 0.58, 0.75, 1.4–2.4, 3.14, 3.3, 5.26, 6.15, 6.47, 6.91, 7.00;

TLC R<sub>f</sub> 0.45 (5% methanol in dichloromethane).

### **EXAMPLE 108**

N-(3-{[6-(3-Bromo-1-ethyl-propyl)-4-hydroxy-2-oxo-2H-65 pyran-3-yl]-cyclopropyl-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula J-3) Refer to Chart J

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To a mixture of 102 mg of the title compound of Preparation 42 and 40  $\mu$ L of pyridine in 1.0 mL of dichloromethane is added 45 mg of 1-methylimidazole-4-sulfonyl chloride. After stirring overnight, the solvent is removed under reduced pressure. The residue is azeotroped with toluene and is then purified by flash column chromatography on silica gel 60 (230–400 mesh) using 2% to 5% methanol in dichloromethane to give 86 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) 80.2, 0.44, 0.60, 0.82, 1.4–2.2, 2.5, 3.1–3.4, 3.62, 5.93, 6.92, 7.07, 7.19, 7.30, 7.40; FAB-MS: [M+H]=550.1037 found;

TLC  $R_f$  0.36 (5% methanol in dichloromethane).

### **EXAMPLE 109**

N-(3-}[6-(3-Azido-1-ethyl-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-cyclopropyl-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula J-4) Refer to Chart J

To a stirring solution of 113 mg of the title compound of Example 108 in 1.2 mL of ethanol is added 50 mg of sodium azide and 0.4 mL of water. The reaction mixture is heated overnight and then cooled. The solvent is removed under reduced pressure and the residue is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 3% to 6% methanol in dichloromethane to give 57 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H-NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ0.25, 0.48, 0.66, 0.90, 1.3–1.8, 2.42, 2.9–3.2, 3.68, 5.94, 6.93, 7.12, 7.19, 7.23, 7.35, 7.46;

FAB-MS: [M+H]=550.1037 found;

TLC R<sub>f</sub> 0.36 (5% methanol in dichloromethane).

# PREPARATION 43

N-(3-{[6-(3-Amino-1-ethyl-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-cyclopropyl-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula J-5) Refer to Chart J

A mixture of 104 mg of the title compound of Example 109 and 30 mg of 5% palladium on charcoal in 2 mL of each of methanol and ethanol is shaken overnight under 45 psi of hydrogen. The mixture is filtered through Celite with methanol washes of the filter cake. The filtrates are combined and the solvent is removed under reduced pressure to give 69 mg of the title compound.

Physical characteristics are as follows:

TLC  $R_f$  0.05 (5% methanol in dichloromethane).

# **EXAMPLE 110**

2-[[8-[[3-[3-[Cyclopropyl[3-[[(1-methyl-1H-imidazol-4-yl) sulfonyl]amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]pentyl]amino]-1,8-dioxooctyl]methylamino]ethane sulfonic acid, monosodium salt (Formula J-6) Refer to Chart J

A suspension of 69 mg of the title compound of Preparation 43 in 1.0 mL of dichloromethane is treated with 0.22 mL (0.65M in acetonitrile) of the triethylamine salt of suleptanic acid and 25  $\mu$ L of diisopropylcarbodiimide. After 1 hour the mixture is treated with 0.5 mL of dimethylformamide. After stirring overnight the solvent is removed under reduced pressure and the residue is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 10% to 30% methanol in dichloromethane. The crude product is dissolved in water saturated n-butanol and partioned with saturated aqueous sodium sulfate. The aqueous phase is extracted twice with additional portions of water saturated n-butanol. The combined n-butanol layers are filtered through a pad of sodium sulfate and concentrated under reduced pressure to give 94 mg of the title compound.

Physical characteristics are as follows:  ${}^{1}\text{H-NMR}$  (CDCl<sub>3</sub>—CD<sub>3</sub>OD)  $\delta 0.05$ –0.6, 0.83, 1.1–2.5, 2.9–3.7, 3.68, 5.84, 6.8–7.6; FAB-MS: [M+H]=786.2838 found; TLC R<sub>4</sub> 0.21 (20% methanol in dichloromethane).

#### EXAMPLES 111-134

Utilizing procedures described above and using starting materials and reagents known and available to one of ordinary skill in organic synthesis, the following additional compounds are prepared:

- 111) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[(2-hydroxy-1,1-bis{hydroxymethyl}-ethyl)-amino]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)
  -benzenesulfonamide
- 112) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[(2-hydroxy-1,1-bis{hydroxymethyl}-ethyl)-amino]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide
- 113) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[(2-hydroxy-1,1-bis{hydroxymethyl}-ethyl)-amino]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide
- 114) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[(2-hydroxy-25 1,1-bis{hydroxymethyl}-ethyl)-amino]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl) -1H-benzoimidazole-2-sulfonamide
- 115) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[(2-hydroxy-1,1-bis{hydroxymethyl}-ethyl)-amino]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide
- 116) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[(2-hydroxy-1,1-bis{hydroxymethyl}-ethyl)-amino]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide
- 117) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{γ-L-glutamyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-benzenesulfonamide
- 118) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{γ-L-glutamyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide
- 119) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{γ-L-glutamyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]- 45 methyl}-phenyl)-1H-imidazole-sulfonamide
- 120) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{γ-L-glutamyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-benzoimidazole-sulfonamide
- 121) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{γ-L-glutamyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide
- 122) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{γ-L-glutamyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide
- 123) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[piperazin-1-yl]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-benzenesulfonamide
- 124) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[piperazin-1-yl]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide
- 125) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[piperazin-1-65 yl]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide

- 126) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[piperazin-1-yl]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-benzoimidazole-2-sulfonamide
- 127) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[piperazin-1-yl]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide
- 128) N-(3-{Cyclopropyl-[4-hydroxy-6-(3-{[piperazin-1-yl]-carbonyl}-amino-propyl)-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide
- 129) 2-[[8-[[3-[3-[Cyclopropyl[3-[[phenylsulfonyl] amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]propyl]amino]-1,8-dioxooctyl]methylamino]-ethanesulfonic acid, monosodium salt
- 130) 2-[[8-[[3-[3-[Cyclopropyl[3-[[(2-pyridyl)sulfonyl] amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]propyl]amino]-1,8-dioxooctyl]methylamino]-ethane sulfonic acid, monosodium salt
- 131) 2-[[8-[[3-[3-[Cyclopropyl[3-[[(1H-benzimidazol-2-yl)sulfonyl]amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]propyl]amino]-1,8-dioxooctyl]methyl amino]-ethanesulfonic acid, monosodium salt
- 132) 2-[[8-[[3-[3-[Cyclopropyl[3-[[(1H-imidazol-2-yl) sulfonyl]amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]propyl]amino]-1,8-dioxooctyl] methylamino]-ethane sulfonic acid, monosodium salt
- 133) 2-[[8-[[3-[3-[Cyclopropyl[3-[[(1-methyl-1H-imidazol-4-yl)sulfonyl]amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]propyl]amino]-1,8-dioxooctyl]methylamino]-ethanesulfonic acid, monosodium salt
- 134) 2-[[8-[[3-[3-[Cyclopropyl[3-[[(1-methyl-1H-imidazol-2-yl)sulfonyl]amino]phenyl]methyl]-4-hydroxy-2-oxo-2H-pyran-6-yl]propyl]amino]-1,8-dioxooctyl]methylamino]-ethane sulfonic acid, monosodium salt

### PREPARATION 44

(Tetrahydropyran-4-yl)-methanol (Formula K-2) Refer to Chart K

To a cold (0°), stirred solution of 651 mg of tetrahydropyran-4-carboxylic acid in 2.5 ml of dry tetrahydrofuran, under argon, is added dropwise 10 ml of a 1.0M solution of borane in tetrahydrofuran. After 18 hours at room temperature, the solution is recooled to 0° and quenched with 1 ml of 1M KOH. The mixture is acidified with 1M aqueous hydrochloric acid and extracted four times with dichloromethane. The extract is dried over magnesium sulfate and concentrated carefully under reduced pressure to afford 0.72 g of the alcohol as a colorless liquid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ1.2–1.4, 1.6, 1.8, 3.3–3.4, 3.6, 4.0 ppm.

### PREPARATION 45

(Tetrahydropyran-4-yl)-methyl p-toluenesulfonate (Formula K-3) Refer to Chart K

To a cold (0°), stirred solution of 5 mmol of the title compound of Preparation 44 and 0.81 ml of pyridine in 5 ml of dichloromethane is added 1.05 g of p-toluenesulfonyl chloride, and the solution is allowed to warm to room temperature. After 18 hours the mixture is partitioned between ethyl acetate and dilute aqueous hydrochloric acid, and the organic phase is washed with brine and dried over magnesium sulfate. Following removal of solvent under reduced pressure, the residue is flash chromatographed on

silica using 50% ethyl acetate in hexane to afford 1.23 g of the title compound as a colorless liquid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ1.2–1.4, 1.6, 1.9–2.0, 2.46, 3.34, 3.85, 3.95, 7.3, 7.8 ppm.

MS: 270

#### PREPARATION 46

(Tetrahydropyran-4-yl)-methyl iodide (Formula K-4) Refer to Chart K

A solution of 800 mg of tosylate of Preparation 45 and 10 887 mg of sodium iodide in 6 ml of acetone is refluxed under nitrogen for six hours, then partitioned between ether and dilute aqueous sodium thiosulfate. The organic phase is washed with brine, dried over magnesium sulfate, and concentrated carefully under atmospheric pressure to give 15 N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-648 mg of the iodide as a colorless liquid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ1.2–1.4, 1.6–1.9, 3.1, 3.37, 3.97 ppm.

### PREPARATION 47

6-(1-(Tetrahydropyran-4-ylmethyl)-propyl)-4-20 hydroxypyran-2-one (Formula K-5) Refer to Chart K

To a cold (-78°) stirred solution of 0.90 ml of disopropylamine in 5 ml of tetrahydrofuran, under argon, is added via syringe 3.7 ml of a 1.6M solution of n-butyllithium in hexane. The solution is warmed to 0°, and after ten minutes, 25 a solution of 431 mg of the title compound of Preparation 50 in 3 ml of hexamethylphosphoramide is added via cannula. After 20 minutes, the deep red solution is cooled to -50°, and 605 mg of iodide of Preparation 46 in 1 ml of tetrahydrofuran is added via cannula. The reaction is allowed to warm slowly to 0° and then quenched by addition of pH 7 30 phosphate buffer. Following removal of tetrahydrofuran under reduced pressure, the residual liquid is acidified with dilute aqueous hydrochloric acid and the resulting precipitate extracted with two portions of ethyl acetate. The organic is washed with dilute aqueous hydrochloric acid and brine, 35 dried over magnesium sulfate, and concentrated under reduced pressure. Flash chromatography of the residue on silica using 5% acetic acid and 30-40% ethyl acetate in dichloromethane provides 553 mg of the title compound as a thick yellow gum.

Physical characteristics are as follows:

TLC R, 0.36 (5% acetic acid, 65% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ0.85, 1.2–1.8, 2.45, 3.34, 3.9, 5.56, 5.94 ppm. MS: 252

# PREPARATION 48

3-[(3-Benzyloxycarbonylaminophenyl)-cyclopropylmethyl] -6-(1-(tetrahydropyran-4-ylmethyl)-propyl-4-hydroxypyran-2-one (Formula K-6) Refer to Chart K

A solution of 549 mg of alkylation product of Preparation 50 47, 970 mg of 3-benzyloxycarbonylaminophenyl cyclopropyl carbinol, and 60 mg of p-toluenesulfonic acid monohydrate in 5 ml of dichloromethane is refluxed through 10 ml of 3 Å sieves for 18 hours. Following removal of solvent under reduced pressure, the residue is flash chromatographed on silica using 25-100% ethyl acetate in dichloromethane to 5% methanol in ethyl acetate, providing 511 mg of the title compound as a tan solid.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.32 (30% ethyl acetate in dichloromethane) <sup>1</sup>H NMR δ0.2, 0.5, 0.7, 0.8, 1.3–1.7, 3.27, 3.42, 3.86, 5.13, 5.96, 7.1-7.4 ppm.

MS: 531

### PREPARATION 49

3-[(3-Aminophenyl)-cyclopropylmethyl]-6-(1-65 (tetrahydropyran-4-ylmethyl)-propyl-4-hydroxy-pyran-2one (Formula K-7) Refer to Chart K

A mixture of 510 mg of the title compound of Preparation 48, 605 mg of ammonium formate, and 100 mg of 5% palladium on carbon in 8 ml of methanol is stirred under argon for three hours, then filtered through diatomaceous earth. The filtrate is concentrated under reduced pressure, and the residue flash chromatographed on silica using 2-4% methanol in dichloromethane to afford 280 mg of the title amine as a white solid.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.33 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.24, 0.42, 0.53, 0.68, 0.84, 1.1–1.7, 2.35, 3.33, 3.6, 3.9, 5.82, 6.5, 6.83, 6.9, 7.11 ppm.

### **EXAMPLE 135**

propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula K-8) Refer

To a stirred solution of 60 mg of the amine of Preparation 49 and 24 µL of pyridine in 0.5 ml of dichloromethane is added 27 mg of 1-methylimidazole-4-sulfonyl chloride. After 18 hours the reaction is flashed on silica using 3-6% methanol in dichloromethane to afford 70 mg of the title compound as a white solid.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.24 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.12, 0.26, 0.45, 0.60, 0.82, 1.1–1.9, 2.3, 3.3, 3.58, 3.9, 6.00, 6.9-7.5 ppm.

HRMS: 541.2238

#### PREPARATION 50

4-Hydroxy-6-propylpyran-2-one (Formula K-9) Refer to

To a cold (-78°), stirred solution of 6.3 ml of diisopropylamine in 40 ml of dry tetrahydrofuran, under argon, is added 27.5 ml of a 1.6M solution of butyllithium in hexane. The solution is brought to 0°, and into this is cannulated a solution of 2.52 g of 4-hydroxy-6-methyl-2-pyrone of formula K-10 in 20 ml of hexamethylphosphoric triamide. The deep red solution is stirred 30 minutes at 0°, then cooled to -45° for the addition of 1.5 ml of ethyl bromide. The solution is warmed to 0° and quenched with 60 ml of 1N aqueous hydrochloric acid. Tetrahydrofuran is removed under reduced pressure and the residue extracted five times with ethyl acetate. The organic phase is washed with brine, 45 dried over magnesium sulfate, and concentrated under reduced pressure. Flash chromatography of the residue on silica gel using 4% acetic acid and 16% ethyl acetate in dichloromethane provides 2.34 g of the title compound as a waxy yellow solid.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.29 (5% acetic acid and 15% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ0.98, 1.6, 2.4, 5.63, 6.05

### PREPARATION 51

4-Hydroxy-6-phenethyl-2H-pyran-2-one (Formula L-2) Refer to Chart L

To a flame-dried flask containing a stirred solution of 0.90 mL of diisopropylamine in 6 mL of anhydrous tetrahydrofuran at -78° C. under an argon atmosphere is added 4.0 mL of a 1.6M solution of n-butyllithium in hexane. The resulting solution is allowed to warm to 0° C. for 20 min, and is then treated via cannula with a solution of 378 mg of commercially available 4-hydroxy-6-methyl-2-pyrone of formula L-1 in 15 mL of tetrahydrofuran. The resulting red, thick slurry is slowly treated with 6.0 mL of distilled hexamethylphosphoramide and allowed to stir for 30 min. The red,

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cloudy solution is then treated with 0.36 mL of benzyl bromide. The reaction quickly becomes a deep orange solution and is allowed to stir at 0° C. for an additional 60 min. The mixture is quenched with excess 1N aqueous hydrochloric acid and the resulting yellow, biphasic mixture 5 is concentrated to remove the tetrahydrofuran. The resulting mixture is partitioned between dichloromethane and water and the acidic aqueous phase is further extracted with additional portions of dichloromethane. The combined organic phase is dried over magnesium sulfate and then 10 concentrated under reduced pressure. The resulting material is diluted with a large volume of diethyl ether and washed with dilute aqueous hydrochloric acid. The ethereal phase is washed with two additional portions of aqueous hydrochloric acid, once with brine, dried over magnesium sulfate, and 15 finally concentrated under reduced pressure. The residue is flash column chromatographed on silica gel 60 (230-400 mesh) eluting with 1% acetic acid and 20% to 40% ethyl acetate in dichloromethane to give 440 mg of the title compound as a tan solid.

Physical characteristics are as follows: <sup>1</sup>H NMR .82.7, 3.0, 5.46, 5.84, 7.1–7.3.

TLC R<sub>f</sub> 0.38 (1% acetic acid and 25% ethyl acetate in dichloromethane.)

MP 137°-138° C.

#### PREPARATION 52

6-(α-Ethyl-phenethyl)-4-hydroxy-2H-pyran-2-one (Formula L-3) Refer to Chart L

To a cold (-78° C.) stirred solution of 0.29 ml of diisopropylamine in 4 ml of dry tetrahydrofuran, under argon, is added 1.2 ml of a 1.6M solution of n-butyllithium in hexane. The solution is warmed to 0° C., kept at that temperature for ten minutes, then cooled to -30° C. Into this solution is cannulated a solution of 189 mg of the title compound of Preparation 51 in 4 ml of tetrahydrofuran. The resulting heterogeneous mixture is warmed to 0°, and sufficient hexamethylphosphoramide (ca 1 ml) is added to render the mixture mostly homogeneous. After the mixture is stirred for 30 minutes at 0° C., 77  $\mu$ L of ethyl iodide is added dropwise. After another 90 minutes, the reaction is quenched with excess 1N aqueous hydrochloric acid, and tetrahydrofuran is removed under reduced pressure. The residue is extracted with three portions of ethyl acetate, and the combined organic extract washed with dilute aqueous hydrochloric acid, dried over magnesium sulfate, and concentrated under reduced pressure. The residue is flash chromatographed on silica gel 60 (230-400 mesh) using 1% acetic acid and 25% ethyl acetate in dichloromethane to provide 182 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.85, 1.6, 2.6, 2.9, 5.59, 5.86, 7.0–7.3.

FAB MS [m+H]=245.1185.

TLC  $R_f$  0.33 (1% acetic acid and 25% ethyl acetate in dichloromethane.)

### PREPARATION 53

 $3-(\alpha-Cyclopropyl-meta-(benzyloxycarbonylamino)benzyl)-6-(\alpha-ethyl-phenethyl)-4-hydroxy-2H-pyran-2-one (Formula L-4) Refer to Chart L$ 

A mixture of 181 mg of the title compound of Preparation 60 52, 220 mg of the compound of formula F-5, 28 mg of p-toluenesulfonic acid monohydrate, and 600 mg of 3 Å molecular sieves in 2 ml of benzene is refluxed under argon for 21 hours, then cooled and filtered through Celite. The filtrate is concentrated under reduced pressure, and the 65 residue flash chromatographed on silica gel 60 (230-400 mesh) using 50-100% ethyl acetate in hexane to provide 250

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mg of a mixture of materials. This is re-subjected to silica gel chromatography, using 5-20% ethyl acetate in dichloromethane, to afford 154 mg (40%) of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.26, 0.48, 0.67, 0.81, 1.6, 1.8, 2.5, 2.7, 2.9, 3.48, 5.14, 5.86, 6.81, 7.0–7.5, 9.46.

EI HRMS m/z=523.2350.

TLC R<sub>f</sub> 0.27 (5% ethyl acetate in dichloromethane.)

#### PREPARATION 54

3-(α-Cyclopropyl-meta-aminobenzyl)-6-(α-ethyl-phenethyl)-4-hydroxy-2H-pyran-2-one (Formula L-5) Refer to Chart L

A mixture of 146 mg of the title compound of Preparation 53 and 50 mg of 5% palladium on carbon in 2 ml of methanol is shaken under 40 psi of hydrogen for two hours, then filtered through Celite. The filtrate is concentrated under reduced pressure to give 105 mg (96%) of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.25, 0.5, 0.65, 0.81, 1.6, 2.5, 2.7, 2.9, 3.4, 5.79, 6.5, 6.8–7.3.

TLC  $R_f$  0.38 (30% ethyl acetate in dichloromethane).

### **EXAMPLES 136-150**

Utilizing procedures analogous to those described above, and reacting the compound of formula L-5 with the appropriate sulfonyl chloride, the following additional compounds of the present invention are prepared. Individual stereoisomers are prepared by chiral HPLC resolution of intermediates such as the compounds of formulas L-3, L-4, L-5 and L-6. (Refer to Chart L).

136) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1methyl-1H-imidazole-4-sulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.29 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.2, 0.5, 0.65, 0.86, 1.63, 1.80, 2.51, 2.8, 3.3, 3.62, 5.7, 6.8–7.4 ppm.

HRMS: 533.1998

137) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.32 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.18, 0.43, 0.63, 0.83, 1.6, 1.75, 2.5, 2.7–2.9, 3.3, 3.55, 5.76, 6.9–7.4 ppm.

HRMS: 533.1983

138) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo- 2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide

Physical characteristics are as follows:

TLC R<sub>e</sub> 0.30 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.2, 0.5, 0.65, 0.86, 1.63, 1.80, 2.51, 2.8, 3.3, 3.62, 5.7, 6.8–7.4 ppm.

HRMS: 533.1993

139) N-(3-(R or S)-{Cyclopropyl-[6-(1-(S)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.30 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.2, 0.5, 0.65, 0.86, 1.63, 1.80, 2.51, 2.8, 3.3, 3.62, 5.7, 6.8–7.4 ppm.

HRMS: 533.1993

140) N-(3-(R or S)-{Cyclopropyl-[6-(1-(S)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-4-sulfonamide

Physical characteristics are as follows:

TLC R<sub>c</sub> 0.30 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.17, 0.44, 0.62, 0.83, 1.6, 1.75, 2.50, 2.7–3.0,

3.3, 3.53, 5.80, 6.9-7.4 ppm.

HRMS: 533.1990

141) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.34 (30% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ0.2, 0.45, 0.6, 0.86, 1.5–1.9, 2.5, 2.8–3.0, 3.2, 5.7, 6.9–7.4, 7.8, 8.6 ppm.

MS: 530

142) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.35 (30% ethyl acetate in dichloromethane) <sup>1</sup>H NMR 80.11, 0.20, 0.43, 0.58, 0.85, 1.5-1.8, 2.5,

2.7–3.0, 3.3, 5.69, 6.9–7.4, 7.8, 8.6 ppm.

MS: 530

143) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide

Physical characteristics are as follows:

TLC  $R_f$  0.34 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.19, 0.5, 0.65, 0.89, 1.6–1.9, 2.5, 2.8–3.0, 3.3, 3.40, 5.70, 6.8–7.4 ppm.

MS: 533

144) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.34 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.20, 0.44, 0.65, 0.88, 1.6–1.8, 2.5, 2.8–3.0, 3.3, 3.42, 5.73, 6.8–7.4 ppm.

MS: 533

145) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1-methyl-1H-imidazole-2-sulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.22 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.16, 0.24, 0.47, 0.64, 0.86, 1.2–1.9, 3.2–3.4,

3.47, 3.7–4.0, 5.89, 6.9–7.4 ppm.

MS: 541

145A) N-(3-[Cyclopropyl[4-hydroxy-2-oxo-6-[1-[ (tetrahydro-2H-pyran-3-yl)methyl]propyl]-2H-pyran-55 3-yl]methyl]phenyl]-8-quinolinesulfonamide

Physical characteristics are as follows:

MW Found: m/z 588.

145B) N-(3-[Cyclopropyl[4-hydroxy-2-oxo-6-[1-[ (tetrahydro-2H-pyran-3-yl)methyl]propyl]-2H-pyran-60 3-yl]methyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide

Physical characteristics are as follows:

MW Found: m/z 541.

146) N-(3-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4- 65 hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-benzimidazole-2-sulfonamide

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Physical characteristics are as follows:

TLC R<sub>f</sub> 0.40 (50% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR 80.1–0.6, 0.85, 1.5–1.7, 2.5, 2.7–3.0, 3.3, 5.74, 6.7–7.3, 7.5–7.7 ppm.

HRMS: 570.2054

147) N-(3-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide

Physical characteristics are as follows:

TLC R, 0.31 (5% methanol in dichloromethane)

<sup>1</sup>H NMR 80.2, 0.4, 0.6, 0.87, 1.5–1.8, 2.5, 2.8–3.0, 3.3, 5.54, 6.8, 6.9–7.4 ppm.

148) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-cyanobenzenesulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.47 (20% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ0.1, 0.2, 0.4, 0.6, 0.84, 1.5–1.8, 2.5, 2.7–3.0,

3.3, 5.70, 6.9, 7.0–7.3, 7.6, 7.8 ppm.

HRMS: 554.1886

149) N-(3-(R or S)-{Cyclopropyl-[6-(1-(R)-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-cyanobenzenesulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.35 (15% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ0.1, 0.2, 0.4, 0.6, 0.85, 1.5–1.9, 2.5, 2.7–3.0,

3.3, 5.7, 6.9–7.3, 7.6, 7.8 ppm.

HRMS: 554.1876

150) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-nitrobenzenesulfonamide

Physical characteristics are as follows:

TLC R, 0.28 (10% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ0.1, 0.2, 0.4, 0.6, 0.83, 1.5–1.9, 2.5, 2.7–3.0, 3.3, 5.70, 6.9–7.3, 7.9, 8.2 ppm.

HRMS: 574.1773

### PREPARATION 55

(2-(2-(2-Methoxyethoxy)-ethoxy)-ethoxy)-ptoluenesulfonate (Formula M-2) Refer to Chart M

To a stirred suspension of 19.1 g of p-toluenesulfonyl chloride in 100 ml of dichloromethane is added a mixture of 16 ml of triethylene glycol monomethyl ether and 10 ml of pyridine, followed by 200 mg of dimethylaminopyridine. After three days the mixture is concentrated under reduced pressure, and the residue partitioned between ethyl acetate and dilute aqueous hydrochloric acid. The organic phase is washed with water, aqueous sodium bicarbonate, and brine, and dried over magnesium sulfate. After removal of solvent under reduced pressure, the residue is flash chromatographed on silica using 25% ethyl acetate in dichloromethane to afford 18.25 g of the title compound as a colorless liquid.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.27 (20% ethyl acetate in dichloromethane) <sup>1</sup>H NMR 82.45, 3.38, 3.5-3.8, 4.15, 7.35, 7.8 ppm. IR 2879, 1357, 1190, 1177, 1108, 1099, 924, 665 cm<sup>-1</sup>

MS: 318

### PREPARATION 56

2-Hydroxy-4-(2-(2-(2-methoxyethoxy)-ethoxy)-acetophenone (Formula M-3) Refer to Chart M

A mixture of 1.52 g of 2,4-dihydroxyacetophenone, 3.82 g of the tosylate of Preparation 55, 3.26 g of cesium carbonate, and 0.2 g of potassium iodide in 20 ml of dioxane is heated overnight at 100°, then cooled and partitioned between dichloromethane and dilute aqueous hydrochloric

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acid. The aqueous phase is extracted with two additional portions of dichloromethane, and the combined organic phase dried over magnesium sulfate and then concentrated under reduced pressure. Flash chromatography of the residue on silica gel using 80-100% ethyl acetate in hexane provides 2.91 g of the title compound as a nearly colorless liquid.

Physical characteristics are as follows: TLC R<sub>f</sub> 0.35 (80% ethyl acetate in hexane) <sup>1</sup>H NMR δ2.56, 3.38, 3.5–3.9, 4.2, 6.4–6.5, 7.6 ppm. IR 1635, 1372, 1257, 1133 cm<sup>-1</sup> MS: 298

#### PREPARATION 57

3-(2-Hydroxy-4-(2-(2-(2-methoxyethoxy)-ethoxy)-ethoxy)phenyl-3-oxopropionic acid ethyl ester (Formula M-4) Refer 15 to Chart M to Chart M

To a stirred solution of 1.49 g of the title compound of Preparation 56 in 10 ml of diethyl carbonate is added, in portions, 600 mg of 60% sodium hydride dispersion in mineral oil. The resulting mixture is heated at 80° for two hours, then cooled and partitioned between dichloromethane and dilute aqueous hydrochloric acid. The organic phase is dried over magnesium sulfate and concentrated under reduced pressure, and the residue purified by flash chromatography on silica gel using 20-30% ethyl acetate in dichloromethane to afford 0.91 g of the title compound as a yellow 25

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.44 (3% acetic acid and 30% ethyl acetate in dichloromethane)

<sup>1</sup>H NMR δ1.3, 3.38, 3.5–4.0, 4.2, 6.4–6.5, 7.6 ppm. MS: 370

#### PREPARATION 58

4-Hydroxy-(2-(2-(2-methoxyethoxy)-ethoxy)-ethoxy)coumarin (Formula M-5) Refer to Chart M

A solution of 789 mg of the title compound of Preparation 35 57 in 10 ml of acetic acid is refluxed for two hours, then concentrated under reduced pressure. Flash chromatography of the residue on silica using 5-10% acetic acid in ethyl acetate provides 634 mg of the title compound as a buff colored solid.

Physical characteristics are as follows: TLC R, 0.31 (10% acetic acid in ethyl acetate) <sup>1</sup>H NMR δ3.37, 3.5–3.9, 4.1, 5.67, 6.6, 6.7, 7.6 ppm. MS: 324

#### PREPARATION 59

3-[(3-Benzyloxycarbonylaminophenyl)-cyclopropylmethyl] -4-hydroxy-7-{2-[2-(2-methoxyethoxy)-ethoxy]-ethoxy}coumarin (Formula M-6) Refer to Chart M

A mixture of 704 mg of the title compound of Preparation 50 58, 775 mg of meta-benzyloxycarbonylaminophenyl cyclopropyl carbinol of formula F-5, and 62 mg of p-toluenesulfonic acid monohydrate in 8 ml of dichloromethane is refluxed for 18 hours through ca. 10 ml of 3 Å sieves. The solution is then concentrated under reduced pressure and the residue flash chromatographed on silica gel using 10-20% of (10% acetic acid in ethyl acetate) in dichloromethane to afford 760 mg of the title compound.

Physical characteristics are as follows:

dichloromethane)

<sup>1</sup>H NMR δ0.27, 0.46, 0.71, 1.61, 3.33, 3.5–3.9, 4.1, 5.13, 6.6, 6.7, 7.1-7.6 ppm.

### PREPARATION 60

3-[(3-Aminophenyl)-cyclopropylmethyl]-4-hydroxy-7-{2- 65 [2-(2-methoxyethoxy)-ethoxy]-ethoxy}-coumarin (Formula M-7) Refer to Chart M

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A solution of 760 mg of the title compound of Preparation 59, 800 mg of ammonium formate, and 200 mg of 5% palladium on charcoal catalyst in 8 ml of methanol is stirred under argon for one hour, then filtered through a pad of diatomaceous earth. The filtrate is concentrated under reduced pressure and the residue triturated with dichloromethane. Removal of solvent under reduced pressure provides 591 mg of the title amine.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.29 (5% methanol in dichloromethane)

### **EXAMPLE 151**

N-(3-{Cyclopropyl-[7-(2-(2-(2-methoxyethoxy)-ethoxy) ethoxy)-4-hydroxycoumarin-3-yl]-methyl}-phenyl)-1methyl-1H-imidazole- 4-sulfonamide (Formula M-8) Refer

To a stirred solution of 70 mg of the title compound of Preparation 60 and 24  $\mu$ L of pyridine in 0.5 ml of dichloromethane is added 27 mg of 1-methylimidazole-4-sulfonyl chloride. After 18 hours, the solution is flash chromatographed on silica gel using 5-15% methanol in dichloromethane, affording 76 mg of the title sulfonamide as a pink amorphous foam.

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.21 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.16, 0.29, 0.45, 0.61, 1.71, 3.34, 3.4–3.9, 4.1, 6.6-6.8, 7.0-7.4, 7.7 ppm.

HRMS: 614.2179

#### EXAMPLES 152-154

Utilizing procedures analogous to those described above, the following additional compounds of the present invention are prepared:

152) N-(3-{Cyclopropyl-[7-methoxy-4hydroxycoumarin-3-yl]-methyl}-phenyl)-1-methyl-

1H-imidazole-4-sulfonamide

Physical characteristics are as follows: TLC R<sub>c</sub> 0.29 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.18, 0.35, 0.50, 0.63, 1.61, 3.51, 3.7, 3.84, 6.7-6.8, 7.1-7.4, 7.7 ppm.

HRMS: 481.1301

153) N-(3-{Cyclopropyl-[7-(2-(2-methoxyethoxy)ethoxy)ethoxy)-4-hydroxycoumarin-3-yl]-methyl}phenyl)-8-quinolinesulfonamide

Physical characteristics are as follows:

TLC R<sub>f</sub> 0.41 (5% methanol in dichloromethane)

<sup>1</sup>H NMR  $\delta$ -0.03, 0.31, 0.47, 1.30, 3.36, 3.5-3.8, 3.9, 4.2, 6.6-7.6, 7.8, 8.0, 8.2 ppm.

HRMS: 661.2219

154) N-(3-{Cyclopropyl-[7-(2-(2-(2-methoxyethoxy)ethoxy)ethoxy)-4-hydroxycoumarin-3-yl]-methyl}phenyl)-2-pyridinesulfonamide

Physical characteristics are as follows:

TLC R<sub>c</sub> 0.31 (5% methanol in dichloromethane)

<sup>1</sup>H NMR δ0.13, 0.34, 0.49, 0.63, 1.6, 3.36, 3.5–3.9, 4.1, 55 6.68, 6.8, 7.1–7.4, 7.6–7.8, 8.5 ppm.

HRMS: 611.2051

### **EXAMPLES 155-190**

The following additional compounds of the present inven-TLC R<sub>f</sub> 0.33 (2% acetic acid and 20% ethyl acetate in 60 tion are prepared by procedures analogous to those described above:

- 155) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]methyl}-phenyl)-2-pyridinesulfonamide
- 156) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]methyl}-phenyl)-4-cyano-2-pyridinesulfonamide

- 157) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-quinolinesulfonamide
- 158) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-5 methyl}-phenyl)-2-hydroxybenzenesulfonamide
- 159) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyrazolesulfonamide
- 160) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-quinazolinesulfonamide
- 161) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-7H-purine-6-sulfonamide
- 162) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide
- 163) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-20 ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-benzimidazole-2-sulfonamide
- 164) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-thiazole-4-sulfonamide
- 165) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-ethoxycarbonyl-1H-imidazole-2-sulfannamide
- 166) N-(3-{Cyclopropyl-[6-(1-(tetrahydropyran-4-ylmethyl)-propyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-3-hydroxy-2-pyridinesulfonamide
- 167) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide
- 168) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-cyano-2-pyridinesulfonamide
- 169) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4- 40 hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-quinolinesulfonamide
- 170) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-hydroxybenzenesulfonamide
- 171) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-pyrazolesulfonamide
- 172) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-2-50 quinazolinesulfonamide
- 173) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-7H-purine-6-sulfonamide
- 174) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide
- 175) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-60 benzimidazole-2-sulfonamide
- 176) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-thiazole-4-sulfonamide
- 177) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-65 hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-4-ethoxycarbonyl-1H-imidazole-2-sulfaonamide

- 178) N-(3-{Cyclopropyl-[6-(1-ethylphenethyl)-4-hydroxy-2-oxo-2H-pyran-3-yl]-methyl}-phenyl)-3-hydroxy-2-pyridinesulfonamide
- 179) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-2-pyridinesulfonamide
- 180) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-4-cyano-2-pyridinesulfonamide
- 181) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-2-quinolinesulfonamide
- 182) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-2-hydroxybenzenesulfonamide
- 183) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-2-pyrazolesulfonamide
- 184) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-2-quinazolinesulfonamide
- 185) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-7H-purine-6-sulfonamide
- 186) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-1H-imidazole-2-sulfonamide
- 187) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-benzimidazole-2-sulfonamide
- 188) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-thiazole-4-sulfonamide
- 189) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-4-ethoxycarbonyl-1H-imidazole-2-sulfaonamide
- 190) N-(3-{Cyclopropyl-[4-hydroxycoumarin-3-yl]-methyl}-phenyl)-3-hydroxy-2-pyridinesulfonamide

### PREPARATION 61

Cyclopropyl-(3-nitrophenyl)methanone (Formula N-2) Refer to Chart N

Charge a jacketed 1 L three neck round bottom flask equipped with stirrer and addition funnel under nitrogen with 580 mL fuming nitric acid and cool to -40° C. Slowly, over 1.5 hours, add cyclopropyl phenyl ketone of formula N-1 (100 g) keeping the temperature below -35° C. Stir 3 hours, monitoring reaction by TLC. Pour reaction mixture into 3 kg ice/water. Extract with 3×500 mL ethyl acetate. Wash combined organic phase with 2×1.5 L saturated aqueous sodium bicarbonate, dry over magnesium sulfate, filter and concentrate to 138 g. Dissolve residue in 270 mL methanol, cool to -20° C. for 18 hours, filter and wash cake with cold methanol. Dry product under reduced pressure for 72 hours, obtaining 63.86 g. GC analysis (15 m. DB-1, T<sub>o</sub>=100° C., 10° C./min., RT -6.0 min.) indicates material to be >98% pure.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ8.86, 8.43, 8.34, 7.70, 2.72, 1.33, 1.17 pm.

IR (Nujol) 2954, 2925, 1664, 1614, 1529, 1442, 1386, 1352, 1225, 1082, 1047, 852, 720, 689 cm<sup>-1</sup>.

Elemental analysis, Found: C, 62.89; H, 4.73; N, 7.32. MS (EI) 191, 150, 104, 69 m/z.

# PREPARATION 62

Cyclopropyl-(3-aminophenyl)methanone (Formula N-3)
Refer to Chart N

Charge platinum on carbon (8.7 g) to Paar bottle. Charge a flask with cyclopropyl(3-nitrophenyl)methanone of Preparation 61 (86.7 g) and methanol (1.56 L) and warm to dissolve, then cool with ice bath to 9° C. Hydrogenate for 50 minutes, keeping temperature below 35° C. and monitoring reaction by TLC. Filter reaction mixture through solka floc, and concentrate under reduced pressure to 70 g.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>2</sub>) δ7.99, 7.47–7.19, 6.84, 3.84, 2.60, 1.23-1.15, 1.03-0.96 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ200.9, 146.8, 139.1, 129.4, 119.3, 118.4, 113.9, 17.2, 11.6 ppm.

### PREPARATION 63

Cyclopropyl-(3-aminocarbobenzoxyphenyl)methanone (Formula N-4) Refer to Chart N

Charge a 3 L round bottom flask equipped with mechanical stirrer and addition funnel under nitrogen with cyclopropyl-(3-aminophenyl)methanone of Preparation 62 (70.0 g), diisopropylethylamine (DIPEA, 90.2 mL) and methylene chloride (CH<sub>2</sub>Cl<sub>2</sub>) (1.3 L). Cool reaction mixture to 0° C. Dilute the benzylchloroformate (67.5 mL) with 15 methylene chloride (186 mL) and add to the substrate solution over one hour keeping temperature at 0°-5° C. A heavy precipitate will form. Allow to warm with stirring for 1.5 hours monitoring reaction by TLC. Pour reaction mixture into 600 mL 1N HCl/600 g ice/4.2 L methylene chloride 20 and stir to dissolve. Separate phases and dry organic phase over magnesium sulfate, filter and concentrate to a dryness. Slurry solids in 3 mL/g hexane, filter, and vacuum dry for

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ8.01, 7.76–7.69, 7.43–7.33, 7.18,

5.21, 2.64, 1.25-1.20, 1.03-0.97 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ200.6, 153.4, 138.7, 138.5, 135.9, 129.3, 128.6, 128.4, 123.1, 122.8, 118.1, 67.2, 17.3, 12.0

### PREPARATION 64

Cyclopropyl-(3-aminocarbobenzoxyphenyl)methanol (Formula N-5) Refer to Chart N

Charge a 2 L three neck round bottom flask equipped with overhead stirrer under nitrogen with cyclopropyl-(3aminocarbobenzoxyphenyl)methanone of Preparation 63 (25 g), tetrahydrofuran (THF) (450 mL) and ethanol (90 mL). Cool reaction mixture to 0°-5° C. and add the sodium borohydride pellets (12.4 g) in three equal portions over 30 minutes. Allow to warm to 23° C. and stir for 20 hours, monitoring reaction by TLC. Recool reaction mixture to 0°-5° C. and slowly quench by adding 90 mL 1N hydrochloric acid, keeping the temperature below 10° C. Pour with stirring into methylene chloride (600 mL) and 1N hydrochloric acid (400 mL). Separate the phases and wash the organic phase with saturated sodium chloride solution (1 L). Dry over magnesium sulfate, filter, and concentrate to

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.41–7.35, 7.33, 7.17, 7.10, 5.17, <sup>50</sup>

3.93, 2.36, 1.16-1.12, 0.60-0.32 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ153.5, 145.0, 137.9, 136.1, 129.0, 128.6, 128.3, 121.2, 117.9, 116.5, 67.9, 67.0, 19.1, 3.6, 2.8 ppm.

### PREPARATION 65

Carbamic acid, [3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-, phenylmethyl ester (Formula N-6) Refer to Chart

A 12-L, three-necked, round-bottomed flask with a Soxhlet extractor containing 3 Å molecular sieves (180 g) and nitrogen inlet is charged with cyclooctene-1-acrylic acid, β, 2-dihydroxy-δ-lactone (59.6 g), p-toluenesulfonic acid (14.9 g), and methylene chloride (7.2 L). The title 65 compound of Preparation 64 (90.0 g) is added, and the reaction mixture is warmed to reflux for 1 h. The reaction

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mixture is then cooled to 20° C. and washed with 1:1 saturated sodium chloride/saturated sodium bicarbonate (3 L), water (3 L), and saturated sodium chloride (3 L), backwashing each aqueous phase with methylene chloride (2x1.5 L). The organic layers are then combined, dried over magnesium sulfate, filtered and concentrated to ca. 1.5 L. The reaction mixture is cooled to -20° C. for 72 h, filtered, and dried under reduced pressure to give 103.5 g. The crude product is then slurried with 12.5 mL/g of hexane, filtered, and dried to give 102.4 g of the title compound. An additional 10.9 g of the title compound is obtained by concentrating the mother liquors from the crystallization and recrystallizing the residue from ethyl acetate.

Physical characteristics are as follows:

MP 113°-115° C. (decomposition).

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.48, 7.38–7.26, 7.17, 6.70, 6.29, 5.20, 3.95, 2.64-2.60, 2.47-2.43, 1.76-1.72, 1.61-1.42,

0.88, 0.73-0.72, 0.63-0.55, 0.29-0.26 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ165.6, 164.0, 161.3, 142.2, 138.5, 129.9, 128.5, 128.3, 128.2, 122.9, 118.0, 117.9, 117.6, 110.7, 106.0, 67.0, 43.7, 30.7, 29.1, 28.8, 26.2, 25.8, 22.1, 13.0, 4.9, 3.8 ppm.

IR (Nujol) 3304, 2995, 2953, 2923, 2855, 1734, 1698, 1665, 1666, 1633, 1610, 1595, 1553, 1491, 1463, 1455, 1445, 1406, 1377, 1313, 1222, 1175, 1085, 1068, 740, 696 25 cm<sup>-1</sup>

MS (EI) m/z 473, 445, 382, 338, 91. For high resolution, Found: 473.2202.

# PREPARATION 66

3-[(3-Aminophenyl)cyclopropylmethyl]-5,6,7,8,9,10hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one (Formula N-7) Refer to Chart N

In a 100-mL, three-necked, round-bottomed flask with a reflux condensor and nitrogen inlet, 10% palladium on carbon (1.0 g) is added to a mixture of the title product of formula N-6, prepared in Preparation 65 (1.95 g) in cyclohexene (50 mL) and the mixture is refluxed for 4 h. The mixture is then filtered through Celite, washed with methylene chloride (CH<sub>2</sub>Cl<sub>2</sub>), and concentrated to give 1.25 g of the title compound as a white solid.

Physical characteristics are as follows:

MP 75°-79° C.

IR (Nujol) 2995, 2951, 2921, 2868, 1660, 1619, 1605, 1590, 1551, 1491, 1460, 1447, 1428, 1404, 1247, 1226, 1202, 1191, 1172, 1126 cm<sup>-1</sup>

MS (EI) m/z 339, 310, 213, 187, 159.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.16, 6.96, 6.84, 6.63, 5.67, 3.87, 2.61, 2.48-2.37, 1.98, 1.75, 1.63-1.26, 0.74-0.65, 0.61-0.53, 0.28-0.22 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ164.2, 161.1, 142.8, 130.2, 117.7, 117.6, 114.7, 114.6, 114.5, 110.9, 106.2, 43.5, 30.6, 29.1, 28.8, 26.2, 25.8, 22.0, 12.8, 4.7, 3.7 ppm.

For high resolution, Found: 339.1845.

# PREPARATION 67

55 4-Cyano-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]benzenesulfonamide (Formula O-3 wherein R<sub>61</sub> is 4-cyanophenyl) Refer to Chart O

A solution of the title product of Preparation 66 (660 mg), pyridine (320 µL), and 4-cyanobenzenesulfonyl chloride (440 mg) in dichloromethane (40 mL) is stirred at room temperature for 18 hr. The crude reaction mixture is evaporated to a volume of 5 ml and chromatographed on silica gel using 50% ethyl acetate in hexane as eluent to give the title compound (641 mg) as a white amorphous solid. This amorphous solid is alternatively crystallized from acetone-:hexane to give 499 mg.

Physical characteristics are as follows:

White solid mp: 183°-183.5° C.

Elemental analysis: found, C, 66.76; H, 5.68; N, 5.38; s,

MS(EI): 504, 476, 463, 338, 309, 233, 220, 207, 195, 186, <sub>5</sub> 153, 144, 130, 117, 102.

HRMS: 504.1710.

TLC(silica gel GF): R<sub>f</sub>=0.4 in 50% ethyl acetate in

#### **EXAMPLE 191**

Disodium-4-cyano-N-[3-[cyclopropyl(5,6,7,8,9,10hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-benzenesulfonamide

To 12.6 g of the title product of Preparation 67 is added 500 ml of methanol and, with rapid stirring, 50 ml of a 1N 15 aqueous NaOH solution. The reaction solution is allowed to stir at room temperature for 1 hour. The yellow solution is evaporated to dryness at 35° C. and the resulting amorphous residue is dissolved in absolute ethanol and re-evaporated to dryness. The yellow residue is kept under high vacuum at 20 methylene chloride. room temperature for 18 hours to yield 14 g of a yellow amorphous solid.

Physical characteristics are as follows:

TLC(silica gel GF): R<sub>=</sub>0.8 streak from the origin (20% ethylacetate in methylene chloride)

K.F. Water: 6.16%

Melt Solvate: 4.2% ethanol

Weight Loss at Room Temperature: 4.99%

Ash: found: 7.83%; Calc'd: 7.50% (corrected for 6.16% water and 4.2% ethanol)

### PREPARATION 68

N-methyl-3[(3-aminophenyl)cyclopropylmethyl]-5,6,7,8,9, 10-hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one

To 678 mg of the title product of Preparation 66 is added 100 ml of absolute ethanol and 330 mg of 10% Pd/C. 183 microliters of a 35% CH<sub>2</sub>O/H<sub>2</sub>O solution is added and the mixture allowed to shake on a Paar apparatus, under 50 lbs of hydrogen, for 2 hours at room temperature. The reaction is filtered over celite and the filter cake is washed well with ethanol. The resulting amber solution is evaporated to dryness. The resulting residue is chromatographed using 10% ethyl acetate in methylene chloride to give 110 mg of the title product. This material is used without further purification in the synthesis of the following sulfonamides.

Physical characteristics are as follows:

TLC(silica gel GF): R<sub>f</sub>=0.5 in 10% ethyl acetate in methylene chloride.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.19, 6.90, 6.71, 6.54–6.52, 3.90, 2.80, 2.63–2.59, 2.43–2.39, 1.75–1.26, 0.70–0.53, 0.28–0.22

### EXAMPLE 192

4-Cyano-N-methyl-N-[3-[cyclopropyl(5,6,7,8,9,10hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-benzenesulfonamide

A solution of the title compound of Preparation 68 (35) mg), pyridine (16  $\mu$ L), and 4-cyanobenzenesulfonyl chloride (20.1 mg) in dichloromethane (2 mL) is stirred at room temperature for 18 hr. The crude reaction mixture is chromatographed on silica gel using 10% ethyl acetate in meth- 60 ylene chloride as eluent to give 27 mg of the title compound as a white amorphous solid.

Physical characteristics are as follows:

MS(EI): 518, 490, 352, 233, 207, 172, 158, 143, 129, 115, 102, 81, 54, 43.

TLC(silica gel GF): R<sub>=</sub>0.7 in 10% ethyl acetate in methylene chloride.

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<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.75–7.72, 7.63–7.60, 7.38–7.19, 6.97–6.94, 6.62, 3.86, 3.19, 2.66–2.62, 2.54–2.50, 1.76–1.20, 0.70–0.59, 0.47–42, 0.24–0.19 ppm.

# **EXAMPLE 193**

4-Fluoro-N-methyl-N-[3-[cyclopropyl(5,6,7,8,9,10hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-benzenesulfonamide

A solution of the title compound of Preparation 68 (20 mg), pyridine (11 µL), and 4-fluorobenzenesulfonyl chloride (10.7 mg) in dichloromethane (2 mL) is stirred at room temperature for 18 hr. The crude reaction mixture is chromatographed on silica gel using 10% ethyl acetate in methylene chloride as eluent to give 19 mg of the title compound as a white amorphous solid.

Physical characteristics are as follows:

MS(EI): 512, 483, 470, 366, 352, 324, 247, 227, 207, 172, 158, 147, 118, 55.

HRMS: Found: 512.1915

TLC(silica gel GF): R<sub>f</sub>=0.7 in 10% ethyl acetate in

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.53–7.48, 7.33–7.23, 7.13–7.07, 6.99-6.97, 6.38, 3.93, 3.16, 2.63-2.61, 2.49-2.46, 1.76-1.25, 0.78-0.61, 0.51-0.45, 0.30-0.17 ppm.

#### **EXAMPLE 194**

N-methyl-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]benzenesulfonamide

A solution of the title compound of Preparation 68 (33.4) mg), pyridine (16  $\mu$ L), and benzenesulfonyl chloride (16.6 mg) in dichloromethane (2 mL) is stirred at room temperature for 18 hr. The crude reaction mixture was chromatographed on silica gel using 10% ethyl acetate in methylene chloride as eluent to give 20 mg of the title compound as a white amorphous solid.

Physical characteristics are as follows:

TLC(silica gel GF): R<sub>f</sub>=0.7 in 10% ethyl acetate in methylene chloride.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.59–7.41, 7.33–7.23, 6.98–6.96, 6.44, 3.90, 3.16, 2.64-2.60, 2.50-2.48, 1.75-1.20, 0.67-0.40, 0.23-0.20 ppm.

# **EXAMPLE 195**

N-methyl-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]-45 1H-Imidazole-1-methyl-sulfonamide

A solution of the title compound of Preparation 68 (33.4 mg), pyridine (16  $\mu$ L), and N-methyl-imidazole-3-sulfonyl chloride (16 mg) in dichloromethane (2 mL) is stirred at room temperature for 18 hr. The crude reaction mixture is chromatographed on silica gel using 50% ethyl acetate in methylene chloride as eluent to give 28 mg of the title compound as a white amorphous solid.

Physical characteristics are as follows:

TLC(silica gel GF): R<sub>f</sub>=0.5 in 50% ethyl acetate in methylene chloride.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.43, 7.33, 7.27–7.15, 3.84–3.81, 3.69, 3.35, 2.63-2.59, 2.50-2.46, 1.75-1.26, 0.68, 0.55, 0.47-0.42, 0.24-0.20 ppm.

Utilizing procedures analogous to those described above, the following compounds of the present invention are prepared:

196) 5-cyano-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-pyridinesulfonamide

197) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-quinolinesulfonamide

- 198) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-imidazolesulfonamide
- 199) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] 5 phenyl]-2-pyrimidinesulfonamide
- 200) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-benzimidazolesulfonamide
- 201) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-quinazolinesulfonamide
- 202) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] <sub>15</sub> phenyl]-6-purinesulfonamide
- 203) 5-cyano-N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-N-methyl-2-pyridinesulfonamide
- 204) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-20 hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-N-methyl-2-quinolinesulfonamide
- 205) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-N-methyl-2-imidazolesulfonamide 25
- 206) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-N-methyl-2-pyrimidinesulfonamide
- 207) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-benzimidazolesulfonamide
- 208) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-2-quinazolinesulfonamide
- 209) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-N-methyl-6-purinesulfonamide
- 210) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] 40 phenyl]-N-methyl-4-thiazolesulfonamide
- 211) N-[3-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl] phenyl]-N-methyl-2-pyridinesulfonamide
- 212) 5-cyano-N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-pyridinesulfonamide
- 213) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-50 quinolinesulfonamide
- 214) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-imidazolesulfonamide
- 215) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'- cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-pyrimidinesulfonamide
- 216) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-benzimidazolesulfonamide
- 217) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-quinazolinesulfonamide
- 218) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'- 65 cyclopropylmethylphenyl)-2-pyrone]-N-methyl-6-purinesulfonamide

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- 219) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-4-thiazolesulfonamide
- 220) N-[6-(1'-benzylpropyl)-4-hydroxy-3-(1'-cyclopropylmethylphenyl)-2-pyrone]-N-methyl-2-pyridinesulfonamide
- 221) 5-cyano-N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2-pyridinesulfonamide
- 222) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2-quinolinesulfonamide
- 223) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2-imidazolesulfonamide
- 224) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2pyrimidinesulfonamide
- 225) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2benzimidazolesulfonamide
- 226) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2quinazolinesulfonamide
- 227) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-6-purinesulfonamide
- 228) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-4-thiazolesulfonamide
- 229) N-[3-(1'-cyclopropylmethylphenyl)-4hydroxycoumarin]-N-methyl-2-pyridinesulfonamide
- 230) 5-cyano-N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-pyridinesulfonamide
- 231) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-quinolinesulfonamide
- 232) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-imidazolesulfonamide
- 233) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-pyrimidinesulfonamide
- 234) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-benzimidazolesulfonamide
- 235) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-ouinazolinesulfonamide
- 236) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-6-purinesulfonamide
- 237) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-4-thiazolesulfonamide
- 238) N-[3-[1-(4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl)-propyl]-phenyl]-N-methyl-2-pyridinesulfonamide

### **EXAMPLE 239**

- 60 2-Pyridylsulfonamide, N-[4-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-(Formula P-2, R is 2-pyridyl) Refer to Chart
  - 3-[(3-Aminophenyl)cyclopropylmethyl]-5,6,7,8,9,10hexahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one of Preparation 66 (100 mg) is dissolved in methylene chloride (3 mL) and pyridine (70  $\mu$ L) added. 2-Pyridylsulfonyl chlo-

ride (52 mg) is added and the solution stirred for 2 hr at 25° C. Chloroform (25 mL) is added and the combined extracts washed with 1N.HCl (20 mL) and dried over sodium sulfate. Removal of the solvent gives a pink gum which is chromatographed over silica gel using the flash column technique eluting with 60% ethyl acetate-hexane. The title compound is obtained as a white solid (80 mg).

Physical characteristics are as follows:

MS m/z 480, 339, 338, 186, 145, 144, 132, 130, 78, 55.

#### **EXAMPLE 240**

4-Pyridylsulfonamide, N-[4-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-(Formula P-2, R is 4-pyridyl) Refer to Chart D

Using procedures described in Example 239, the title compound is obtained as a white solid.

Physical characteristics are as follows:

MS m/z 480, 338, 207, 186, 145, 144, 117, 79, 78, 55

### **EXAMPLE 241**

5-Cyanopyridin-2-yl-sulfonamide, N-[4-[cyclopropyl(5,6,7, 8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]-(Formula P-2, R is 5-cyanopyridin-2-yl) Refer to Chart P

The title compound is prepared using procedures described in Example 239.

#### **EXAMPLE 242**

2-Pyrazinylsulfonamide, N-[4-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-(Formula P-2, R is 2-pyrazinyl) Refer to Chart P

The title compound is prepared using procedures described in Example 239.

### EXAMPLE 243

2-Pyrimidinylsulfonamide, N-[4-[cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) methyl]phenyl]-(Formula P-2, R is 2-pyrimidinyl) Refer to Chart P

The title compound is prepared using procedures described in Example 239.

# EXAMPLE 244

4-6-Dimethylpyrimidin-2-yl-sulfonamide, N-[4- <sup>45</sup> [cyclopropyl(5,6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl)methyl]phenyl]-(Formula P-2, R is 4,6-dimethylpyrimidin-2-yl) Refer to Chart P

The title compound is prepared using procedures described in Example 239.

### **EXAMPLE 245**

4-Methylpyrimidin-2-yl-sulfonamide, N-[4-[cyclopropyl(5, 6,7,8,9,10-hexahydro-4-hydroxy-2-oxo-2H-cycloocta[b] pyran-3-yl)methyl]phenyl]-(Formula P-2, R is 55 4-methylpyrimidin-2-yl) Refer to Chart P

The title compound is prepared using procedures described in Example 239.

### PREPARATION 69

6,6-Bis-(2-cyclopropyl-ethyl)-dihydro-pyran-2,4-dione (Formula Q-2) Refer to Chart Q

To a suspension of 150 mg of sodium hydride (60% dispersion in mineral oil) in 4 ml of dry THF under argon atmosphere at 0° C. is added dropwise 0.38 ml of methyl acetoacetate. After 10 minutes 2.3 ml of butyllithium (1.6M in hexanes) is added. After 10 minutes a solution of 0.48 g

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of the compound of formula Q-1 (prepared as described in Preparation 79 (Formula S-4, refer to Chart S)) in 3 ml of tetrahydrofuran is added. The reaction mixture is stirred for 1 hour, then partitioned between ethyl acetate and dilute aqueous hydrogen chloride. The aqueous phase is extracted with two additional portions of ethyl acetate. The organic phases are combined, washed with brine, dried over magnesium sulfate and concentrated under reduced pressure. The residue is diluted with 5 mL of methanol and the 10 resulting solution treated with 12 mL of water followed by 3.0 ml of 1M aqueous sodium hydroxide. After 2 hours of vigorous stirring the methanol is removed under reduced pressure. The aqueous phase is washed once with diethyl ether; the ether phase is discarded. The aqueous phase is cooled to 0° C., then acidified with dilute aqueous hydrogen chloride. The resulting precipitate is extracted with four portions of dichloromethane. The combined dichloromethane extracts are dried over magnesium sulfate and concentrated under reduced pressure. The residue is dis-20 solved in diethyl ether-hexane and the solution is chilled to provide to provide 0.42 g of the title compound as a pale vellow solid.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.0, 0.4, 0.6, 1.2, 1.7, 2.6, 3.4.

#### PREPARATION 70

6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-3-[1-(3-nitrophenyl)-propyl]-pyran-2-one (Formula Q-3) Refer to Chart O

To a stirred solution of 0.41 g of the title compound of Preparation 69 (Formula Q-2) and 0.25 g of the 3-nitrobenzaldehyde in 5 ml of dry tetrahydrofuran is added a solution of 0.44 g of aluminum trichloride in 4.5 ml of tetrahydrofuran. After 2 hours, the reaction mixture is treated with 1.0 g of sodium carbonate decahydrate, stirred 10 minutes, diluted with diethyl ether and finally charged with magnesium sulfate. The resulting mixture is filtered through a pad of Celite with diethyl ether rinses. The filtrates are combined and concentrated under reduced pressure. The resulting residue is charged with 103 mg of copper (I) bromide-dimethyl sulfide complex and 5 ml of dry tetrahydrofuran under an argon atmosphere. The reaction mixture is treated dropwise with 2.5 mL of triethyl aluminum (1.0M in hexane) over 1.5 hours. The reaction is then slowly treated with ice and partitioned between diethyl ether and dilute aqueous hydrogen chloride. The aqueous phase is extracted with three additional portions of diethyl ether. The combined ether extracts are washed with brine, dried over magnesium sulfate, and concentrated under reduced pressure. Flash column chromatography of the residue on silica gel using 20% to 40% ethyl acetate in hexane affords 0.44 g of the title compound as a tan foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.0, 0.4, 0.6, 1.0, 1.2, 1.7–1.9, 2.0–2.4, 2.6, 4.2, 7.5, 7.8, 8.1, 8.3

### PREPARATION 71

3-[1-(3-Amino-phenyl)-propyl]-6,6-bis-(2-cyclopropylethyl)-5,6-dihydro-4-hydroxy-pyran-2-one (Formula Q-4) Refer to Chart Q

To a solution of 0.44 g of the title compound of Preparation 70 (Formula Q-3) in 6 ml of methanol is added 0.65 g of ammonium formate and 50 mg of 10% palladium on carbon. The black slurry is stirred under argon for 3 hours, then filtered through pad of Celite with methanol washes. The filtrates are combined and the solvent is removed under reduced pressure. The residue is triturated with four portions of dichloromethane. The combined dichloromethane washes

are concentrated under reduced pressure to provide  $0.37\,\mathrm{g}$  of the title compound as a white foam.

Physical characteristics are as follows: R<sub>f</sub> 0.08 (50% diethyl ether in hexane)

#### **EXAMPLE 246**

N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula Q-5:  $R_1$  is 1-methylimidazol-4-yl) Refer to Chart Q

To a flask containing 57 mg of the title compound of Preparation 71 (Formula Q-4) and 24  $\mu$ l of pyridine in 1.0 ml of dichloromethane is added 27 mg of 1-methylimidazole-4-sulfonyl chloride. After 6 hours the reaction mixture is concentrated under reduced pressure. The pyridine is azeo-troped thrice with toluene. The resulting residue is flash column chromatographed on silica gel using 2% to 6% methanol in dichloromethane to provide 51 mg of the title compound as a white foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.0, 0.4, 0.6, 0.9, 1.1–1.4, 1.7–2.2, 2.5, 3.7, 3.95, 6.9, 7.1, 7.4, 7.5

HRMS: 528.2537 (FAB)

#### **EXAMPLE 247**

N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide (Formula Q-5:  $R_1$  is 5-cyano-2-pyridyl) Refer to Chart Q

Using the general sulfonylation procedure described in Example 246, 57 mg of the amine of Preparation 71 (Formula Q-4) is reacted with 30 mg of 5-cyanopyridine-2-sulfonyl chloride. Flash column chromatography on silica gel using 1% to 3% methanol in dichloromethane provides 62 mg of the title compound as a tan foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.0, 0.4, 0.6, 0.9, 1.1–1.4, 1.6–2.2, 2.5, 3.95, 6.9–7.2, 8.0, 8.2, 9.0

HRMS: 550.2370 (FAB)

#### PREPARATION 72

3-Aminopropiophenone (Formula R-2) Refer to Chart R

To a solution of 3-nitropropiophenone (Formula R-1) (1.79 g) in diethyl ether is added 5% Pt/C catalyst (0.20 g). The resulting suspension is placed under a hydrogen gas atmosphere and stirred for 6 hours. The reaction mixture is filtered through a pad of Celite and the pad washed with additional portions of diethyl ether. The combined filtrates are concentrated under reduced pressure to provide 1.49 g of the title compound as pale yellow, low melting solid.

Physical characteristics are as follows: <sup>1</sup>H NMR δ1.2, 3.0, 6.9, 7.2–7.4 R<sub>f</sub> 0.45 (33% ethyl acetate in hexane)

# PREPARATION 73

1-[3-(Dibenzyl-amino)-phenyl]-propan-1-one (Formula 55 R-3) Refer to Chart R

To a solution of the title compound of Preparation 72 of Formula R-2 (1.5 g) in dichloromethane (50 mL) is added disopropylethylamine (6.0 mL) followed by benzyl bromide (3.6 mL). After stirring for 6 hours the reaction mixture is heated to reflux overnight. The reaction mixture is cooled to room temperature, diluted with diethyl ether (50 mL) and washed sequentially with dilute aqueous potassium hydrogen sulfate, water, saturated aqueous sodium bicarbonate, and brine. The organic layer is dried over sodium sulfate and 65 concentrated under reduced pressure. The residue is purified by flash column chromatography on silica gel eluting with

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5% to 20% ethyl acetate in hexane to provide 2.38 g of the title compound as pale yellow solid.

Physical characteristics are as follows: 
<sup>1</sup>H NMR δ1.1, 2.9, 4.7, 6.9, 7.2–7.4
Anal. Found: C, 83.88; H, 7.03; N, 4.20
MS: 329 (EI)

### PREPARATION 74

6-[3-(Dibenzyl-amino)-phenyl]-dihydro-pyran-2,4-dione (Formula R-4) Refer to Chart R

Using the general procedure described in Preparation 69 for the formation of the dihyropyranone ring, the compound of Formula R-3 of Preparation 73 (1.96 g) is reacted with the dianion of methyl acetoacetate and cyclized to provide 0.76 g of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.8, 1.9, 2.6–2.9, 3.1–3.2, 4.7, 6.5–6.7, 7.1–7.4 MS: 413 (EI)

### PREPARATION 75

6-[3-(Dibenzyl-amino)-phenyl]-5,6-dihydro-6-ethyl-4-hydroxy-3-[1-(3-nitro-phenyl)-propyl]-pyran-2-one (Formula R-5) Refer to Chart R

Using the general procedure described in Preparation 70, aluminum trichloride catalyzed condensation of 3-nitrobenzaldehyde with the compound of Formula R-4 of Preparation 74 (727 mg), followed by copper catalyzed conjugate addition with triethyl aluminum provides 800 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.6, 1.6–2.1, 2.8, 3.4, 3.8, 4.4, 6.4–6.6, 6.8–7.4, 7.7–8.0

MS: 576 (EI)

#### PREPARATION 76

6-(3-Amino-phenyl)-3-[1-(3-amino-phenyl)-propyl]-6-ethyl-5,6-dihydro-4-hydroxy-pyran-2-one (Formula R-6) Refer to Chart R

Using the general procedure described in Preparation 71, catalytic hydrogenation of the compound of Formula R-5 of Preparation 75 (114 mg) with ammonium formate and Pd/C affords 61 mg of the title compound. Alternatively, the compound of Formula R-5 of Preparation 75 (114 mg) is reduced with Pd/C and hydrogen gas to give 72 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.6–0.9, 1.8–2.1, 3.0, 3.8, 6.4–6.6, 6.95, 7.1 R<sub>s</sub> 0.40 (10% methanol in dichloromethane)

#### **EXAMPLE 248**

N-(3-[1-(6-Ethyl-5,6-dihydro-4-hydroxy-6-[3-([(1-methyl-1H-imidazol- 4-yl)sulfonyl]amino)phenyl]-2-2H-pyran-3-yl)propyl]phenyl)-1-methyl-1H-imidazole-4-sulfonamide (Formula R-7: R<sub>1</sub> is 1-methylimidazol-4-yl) Refer to Chart P

Using the general sulfonylation procedure described in Example 246, the compound of Formula R-6 of Preparation 76 (61 mg) is reacted with 1-methylimidazole-4-sulfonyl chloride to provide 59 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.3–0.7, 1.6–2.0, 3.0, 3.4–3.7, 6.7–7.5 HRMS: 655.1995 (FAB)

### **EXAMPLE 249**

5-Cyano-N-(3-[1-(6-[3-([(5-cyano-2-pyridinyl)sulfonyl] amino)phenyl]-6-ethyl-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl)propyl]phenyl)-2-pyridinesulfonamide (Formula R-7: R is 5-cyano-2-pyridyl) Refer to Chart R

Using the general sulfonylation procedure described in Example 246, the compound of Formula R-6 of Preparation 76 (66 mg) is reacted with 5-cyano-2-pyridine sulfonyl chloride to provide 40 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.3–0.9, 1.3, 1.6–1.9, 3.0, 3.7, 6.6–7.2, 7.9–8.2, 8.8–9.0

HRMS: 699.1679 (FAB)

#### PREPARATION 77

N-Methoxy-N-methyl-4-pentenoic amide (Formula S-2) Refer to Chart S

To a suspension of 4-pentenoic acid (Formula S-1) (2.00 g) and N,O-dimethylhydroxylamine hydrochloride (2.15 g) in dichloromethane (50 mL) at 0° C. is added diisopropylethylamine (11.5 mL) followed by bis(2-0x0-3-0xazolidinyl)phosphinic chloride (5.60 g). After stirring overnight, the reaction mixture is concentrated under reduced pressure. The residue is partioned between dilute aqueous potassium hydrogen sulfate and diethyl ether. The aqueous phase is extracted with two additional portions of diethyl ether. The organic extracts are combined, washed with brine, dried over sodium sulfate and concentrated under reduced pressure. The residue is purified by flash column chromatography on silica gel eluting with 50% to 80% diethyl ether in hexane to provide 2.58 g of the title compound as a tan oil.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 2.3–2.6, 3.20, 3.70, 4.9–5.1, 5.75–5.95  $R_{f}$  0.17 (25% diethyl ether in hexane)

### PREPARATION 78

Nona-1,8-dien-5-one (Formula S-3) Refer to Chart S

To a flame-dried flask under an argon atmosphere containing a solution of the title compound of Preparation 77 (Formula S-2) (1.45 g) in dry tetrahydrofuran (10 mL) at 0° 35 C. is added 3-butenyl-1-magnesium bromide (20 mL, 1M solution in tetrahydrofuran. (Preparation of this Grignard reagent from magnesium metal and 4-bromo-1-butene is described in J.Org.Chem. 43:4247 (1978)). After 1 hour at 0° C., the reaction mixture is warmed to room temperature; after 1 hour at room temperature, the reaction mixture is poured into dilute aqueous potassium hydrogen sulfate and partioned against diethyl ether. The aqueous phase is extracted with three additional portions of diethyl ether. The organic extracts are combined, washed with brine, dried over sodium sulfate and carefully concentrated under reduced pressure. The resulting liquid is purified by distillation to provide 1.32 g of the title compound as a tan oil.

Physical characteristics are as follows: <sup>1</sup>H NMR δ2.3, 2.5, 5.0, 5.7-5.9 R<sub>f</sub> 0.66 (25% diethyl ether in hexane)

### PREPARATION 79

1,5-Dicyclopropyl-pentan-3-one (Formula S-4) Refer to Chart S

To a flame-dried flask under an argon atmosphere equipped with a reflux condenser containing zinc metal (8.0 g) and cuprous chloride (1.25 g) is added a solution of the title compound of Preparation 78 (Formula S-3) (1.32 g) in dry diethyl ether (10 mL). The resulting suspension is 60 charged with diiodomethane (5.0 mL) and the reaction flask placed in 40° C. ultrasound bath (Branson 2200) and sonicated. After 2 hours heating is ceased and sonication is continued overnight. The reaction mixture is then diluted with diethyl ether (50 mL), cooled to 0° C., and treated with excess saturated aqueous ammonium chloride. After 0.25 hours of vigorous stirring, the mixture is filtered and the

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layers separated. The aqueous phase is extracted with two additional portions of diethyl ether. The organic extracts are combined and washed sequentially with dilute aqueous sodium thiosulfate, saturated aqueous sodium bicarbonate, brine; dried over magnesium sulfate and then carefully concentrated under reduced pressure. The resulting residue is purified by flash column chromatography on silica gel eluting with 5% to 20% diethyl ether in hexane to provide 0.48 g of the title compound as an oil.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.0, 0.4, 0.65, 1.45, 2.50 R<sub>f</sub> 0.44 (10% diethyl ether in hexane)

### PREPARATION 80

3-[2,2-Dimethyl-1-(3-nitro-phenyl)-propyl]-5,6-dihydro-4hydroxy-6-phenethyl-6-propyl-pyran-2-one (Formula T-3) Refer to Chart T

To a flame-dried flask containing a slurry of 977 mg of activated zinc metal in 1.0 mL of dry tetrahydrofuran under an argon atmosphere is added 40  $\mu$ L of 1,2-dibromoethane. The mixture is placed in 45° C. ultrasound bath (Branson 2200) and sonicated with stirring. After 10 minutes the mixture is treated with 0.25 mL of chlorotrimethylsilane (1.0M in tetrahydrofuran). After 10 minutes, the mixture is diluted with 4 mL of tetrahydrofuran and treated dropwise with 1.50 mL of 2-iodo-2-methylpropane. The mixture is stirred and sonicated at 45° C. for an additional 3 hours, then cooled to room temperature without stirring. In a separate flask 954 mg of anhydrous lithium chloride is heated in an 110° C. oil bath in vacuo for 1 hour. The LiCl flask is cooled to room temperature, placed under an argon atmosphere and 30 charged with 1.01 g of copper (I) cyanide followed by 10 mL of tetrahydrofuran. After 15 minutes of stirring at room temperature, the LiCl—CuCN mixture is cooled to -30° C. and treated via cannula with the organozinc mixture prepared as described above in the first flask. The reaction flask is warmed from -30° C. to 0° C., stirred 10 minutes then cooled to -78° C. The preparation of this organometallic reagent is analogous to literature procedures (Org. Syn. 70:195-203 (1991)) described for related reagents.

In a separate flask a stirred solution of 1.56 g of 40 6-phenethyl-6-propyldihydro-pyran-2,4-dione of Formula T-2 (prepared from the compound of Formula T-1 as described in Preparation 17 above) and 915 mg of the 3-nitrobenzaldehyde in 22 mL of dry tetrahydrofuran is treated with a solution of 1.60 g of aluminum trichloride in 14 mL of tetrahydrofuran. After 2 hours, the reaction mixture is treated with 3.6 g of sodium carbonate decahydrate, stirred 5 minutes, diluted with diethyl ether and finally charged with magnesium sulfate. The resulting mixture is filtered through a pad of Celite with diethyl ether washes. The filtrates are combined and concentrated under reduced pressure. The resulting residue is charged with 9 mL of dry tetrahydrofuran under an argon atmosphere and is added via cannula to the cooled (-78° C.) organometallic reagent solution prepared as described above. After 0.5 hours the reaction mixture is warmed to 0° C. After 0.5 hours at 0° C. the reaction is poured into cold dilute ammonium chloride and the aqueous phase is made acidic with dilute aqueous hydrogen chloride. The mixture is treated with ethyl acetate and filtered through a pad of Celite with ethyl acetate washes. The layers are separated and the aqueous phase is extracted with three additional portions of ethyl acetate. The combined ethyl acetate extracts are washed with aqueous sodium thiosulfate, brine; dried over magnesium sulfate, and concentrated under reduced pressure. Flash column chromatography of the residue on silica gel eluting with 30% to 50% ethyl acetate in hexane affords 1.73 g of the title compound as a tan foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.9, 1.1, 1.3, 1.6–2.0, 2.5–2.8, 4.3, 6.9–7.3, 7.8, 8.0, 8.5

HRMS: 452.2449 (FAB)

#### PREPARATION 81

3-[1-(3-Amino-phenyl)-2,2-dimethyl-propyl]-5,6-dihydro-4-hydroxy-6-phenethyl-6-propyl-pyran-2-one (Formula T-4) Refer to Chart T

To a solution of 1.72 g of the title compound of Preparation 80 (Formula T-3) in 25 mL of methanol is added 3.0 10 g of ammonium formate and 400 mg of 10% palladium on carbon. The black slurry is stirred under nitrogen for 3 hours, then filtered through pad of Celite with methanol washes. The filtrates are combined and the solvent is removed under reduced pressure. The residue is repeatedly triturated with portions of dichloromethane and the combined dichloromethane washes concentrated under reduced pressure. The residue is flash column chromatographed on silica gel eluting with 10% ethyl acetate in dichloromethane to provide 1.48 g of the title compound as a white foam.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.7–0.9, 1.1, 1.3–2.6, 4.2, 6.55, 6.9–7.3

HRMS: 422.2686 (FAB)

## **EXAMPLE 250**

N-[3-(1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula T-5:  $R_1$  is 1-methylimidazol-4-yl) Refer to Chart T

To a solution of 1.48 g of the title compound of Preparation 81 (Formula T-4) in 25 ml of dichloromethane at 0° C. is added 0.57 mL of pyridine followed by 632 mg of 1-methylimidazole-4-sulfonyl chloride. After 3 hours the reaction mixture is warmed to room temperature and concentrated under reduced pressure. Pyridine is azeotroped thrice with toluene. The resulting residue is flash column chromatographed on silica gel using 2% to 6% methanol in dichloromethane to provide 1.7 g of the title compound as a white solid

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 0.97, 1.35, 1.6–2.0, 2.5–2.7, 3.6, 4.1, 6.9–7.5

HRMS: 566.2684

The individual stereoisomers of this compound are the following:

N-[3-(1(S)-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula KK-8 wherein  $R_4$  is 1-methyl-1H-imidazol-4-yl) Refer to Chart KK;

N-[3-(1(R)-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula LL-8 wherein  $R_4$  is 1-methyl-1H-imidazol-4-yl) Refer to Chart LL;

N-[3-(1(S)-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula MM-8 wherein  $R_4$  is 1-methyl-1H-imidazol-4-yl) Refer to Chart MM; and

N-[3-(1(R)-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula NN-8 wherein R<sub>4</sub> is 1-methyl-1H-imidazol-4-yl) Refer to Chart NN.

#### **EXAMPLE 251**

5-Cyano-N-[3-(1-[5,6-dihydro-4-hydoxy-2-oxo-6-(2-phenethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)

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phenyl]-2-pyridinesulfonamide (Formula T-5: R<sub>1</sub> is 5-cyano-2-pyridyl) Refer to Chart T

Using the general sulfonylation procedure described in Example 250, 42 mg of the amine of Preparation 81 (Formula T-4) is reacted with 20 mg of 5-cyanopyridine-2-sulfonyl chloride. Flash column chromatography on silica gel using 1% to 3% methanol in dichloromethane provides 56 mg of the title compound as a white foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 0.92, 1.35, 1.6–2.0, 2.5–2.7, 4.0, 6.9–7.4, 8.0, 8.9

HRMS: 588.2532

#### PREPARATION 82

N-Methoxy, N-methyl 3-(4-fluorophenyl)propionamide (Formula U-2) Refer to Chart U

To a cold (0°), stirred solution of 5.0 g of 3-(4-fluorophenyl)propionic acid of Formula U-1, 3.2 g of (N,O)-dimethylhydroxylamine hydrochloride, and 11.4 ml of diisopropylethylamine in 40 ml of dichloromethane is slowly added a solution of 5.0 ml of diethyl cyanophosphonate in 10 ml of dichloromethane. After 18 hours, the solution is concentrated under reduced pressure. The residue is dissolved in ethyl acetate, and the solution washed with dilute HCl, water, aqueous sodium bicarbonate, and brine, and dried over magnesium sulfate. Removal of the solvent under reduced pressure provides 6.94 g of the title compound.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 2.7, 2.9, 3.17, 3.61, 7.0, 7.2 ppm IR 1665, 1511, 1222, 1033, 990 cm $^{-1}$  TLC  $R_f$  0.34 (5% ethyl acetate in dichloromethane)

### PREPARATION 83

1-(4-Fluorophenyl)-3-hexanone (Formula U-3) Refer to Chart U

A stirred solution of 4.68 g of the title compound of Preparation 82 (Formula U-2) in 25 ml of dry THF under argon is cooled to -15°, and to the solution is added 17 ml of a 1M solution of propylmagnesium chloride in ether. The resulting solid mass is warmed to 0°, kept at that temperature for 90 minutes, then partitioned between ether and cold dilute HCl. The aqueous phase is extracted with one additional portion of ether, and the combined organic phase washed with brine and dried over magnesium sulfate. Following removal of solvent by distillation at atmospheric pressure, the residue is purified by evaporative distillation (ca 160° @ 13 mmHg) to provide 3.51 g of the title compound as a colorless liquid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.89, 1.6, 2.36, 2.7, 2.9, 6.9, 7.1 ppm IR 2965, 1714, 1511, 1222 cm<sup>-1</sup>

## PREPARATION 84

5,6-Dihydro-4-hydroxy-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-2-one (Formula U-4) Refer to Chart U

To a cold (0°), stirred slurry of 950 mg of sodium hydride (60% dispersion in mineral oil) in 30 ml of dry THF, under argon, is added dropwise 2.3 ml of methyl acetoacetate. After 5 minutes, 13.5 ml of butyllithium (1.6M in hexanes) is added, and the mixture stirred another 5 minutes before addition of a solution of 3.51 g of the title compound of Preparation 83 (Formula U-3) in 4 ml of THF. The solution is stirred for 1 hour, then partitioned between ethyl acetate and cold dilute HCl. The aqueous phase is extracted with two additional portions of ethyl acetate, and the combined organic phase washed with brine and dried over magnesium sulfate. Removal of the solvent under reduced pressure

provides the intermediate ester with the following physical characteristics: TLC R, 0.45 (50% ethyl acetate in hexane).

The ester is stirred in 20 ml of 1M sodium hydroxide, 80 ml of water, and 40 ml of methanol for 90 minutes, then the methanol is removed under reduced pressure. The aqueous phase is washed once with ether, the ether phase being discarded, and then acidified with dilute HCl. The resulting precipitate is extracted with four portions of dichloromethane, and the extract dried over magnesium sulfate and concentrated under reduced pressure. The residue is dissolved in 1:1 ether-hexane and the solution chilled to provide crystals, which are filtered, washed with ether-hexane, and dried under vacuum to afford 3.24 g of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.96, 1.4, 1.8, 2.0, 2.5, 2.7, 7.0, 7.1 ppm
IR 2962, 1655, 1604, 1510, 1221 cm<sup>-1</sup>
M.P. 113°-114.5°
Anal. Found: C, 68.85; H, 6.99
MS: M+ 278
R<sub>f</sub> 0.44 (5% methanol in dichloromethane)

#### PREPARATION 85

3-(1-(3-Benzyloxycarbonylaminophenyl)-2,2-dimethylpropyl)-5,6-dihydro-4-hydroxy-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-2-one (Formula U-6: R<sub>1</sub> is tert-butyl) Refer to Chart U

To a stirred solution of 3.06 g of the title compound of Preparation 84 (Formula U-4) and 2.81 of the title compound of Preparation 6 above (Formula B-2) in 30 ml of dry THF is added a solution of 2.93 g of AlCl<sub>3</sub> in 20 ml of THF. After two hours, 6.4 g of sodium carbonate decahydrate is added, and after five minutes the mixture is filtered through Celite with ether rinses. Removal of the solvent under reduced pressure provides the intermediate benzylidene compound of Formula U-5.

To this is added, under argon, 1.13 g of copper (I) bromide-dimethyl sulfide complex and 30 ml of THF, and the mixture is cooled to 0° for dropwise addition of 18.1 ml of tert-butylmagnesium chloride (1.0M in THF). After 10 minutes, the reaction is partitioned between ether and cold dilute HCl. The organic phase is washed with brine, dried over magnesium sulfate, and concentrated under reduced pressure. Flash chromatography of the residue on silica gel using 30–35% ethyl acetate in hexane affords 1.83 g of the title compound as a foam.

Physical characteristics are as follows: <sup>1</sup>H NMR 80.87, 1.1, 1.3, 1.6–2.2, 2.5, 5.12, 6.8–7.6 ppm HRMS: 574.2955

R<sub>f</sub> 0.29 (35% ethyl acetate in hexane)

### PREPARATION 86

3-(1-(3-Aminophenyl)-2,2-dimethylpropyl)-5,6-dihydro-4-hydroxy-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-2-one (Formula U-7:  $R_1$  is tert-butyl) Refer to Chart U

A mixture of 1.83 g of the title compound of Preparation 85 (Formula U-6), 2.0 g of ammonium formate, and 400 mg 55 of 10% palladium on carbon in 25 ml of methanol is stirred under argon for 90 minutes, then filtered through Celite. The solvent is removed under reduced pressure and the residue flash chromatographed on silica gel using 10% ethyl acetate in dichloromethane to provide 1.24 g of the title compound 60 as a white foam.

Physical characteristics are as follows:

R<sub>c</sub> 0.28 (10% ethyl acetate in dichloromethane)

The compound of Formula U-7 wherein  $R_1$  is ethyl is prepared from U-4 by analogous procedures as in the preparation of U-7 wherein  $R_1$  is tert-butyl (Preparations 85 and 86).

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#### **EXAMPLE 252**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula U-8:  $R_1$  is tert-butyl,  $R_2$  is 1-methylimidazole-4-yl) Refer to Chart U

To a cold (0°), stirred solution of 88 mg of the title compound of Preparation 86 (Formula U-7) and 32  $\mu$ l of pyridine in 0.5 ml of dichloromethane is added 36 mg of 1-methylimidazole-4-sulfonyl chloride. After 90 minutes the reaction mixture is flash chromatographed on silica using 3-4% methanol in dichloromethane to provide 112 mg of the title compound as a white foam.

Physical characteristics are as follows:

5 <sup>1</sup>H NMR δ0.8–1.0, 0.96, 1.3, 1.7, 2.35, 2.5, 3.6, 3.7, 6.8–7.5 ppm

HRMS: 583.2525

 $R_f$  0.31 (5% methanol in dichloromethane)

#### **EXAMPLE 253**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula U-8:  $R_1$  is tert-butyl,  $R_2$  is 5-cyanopyridine-2-yl) Refer to Chart U

Using the general sulfonylation procedure of Example 252, 88 mg of the amine of Preparation 86 (Formula U-7,  $R_1$  is tert-butyl) is reacted with 5-cyanopyridine-2-sulfonyl chloride. Flash chromatography on silica using 10–15% ethyl acetate in dichloromethane provides 107 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows: <sup>1</sup>H NMR 80.92, 1.3, 1.7, 2.5, 6.8–7.5, 8.0, 8.9 ppm HRMS: 606.2423

### **EXAMPLE 254**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula U-8:  $R_1$  is ethyl,  $R_2$  is 1-methylimidazole-4-yl) Refer to Chart U

Using the general sulfonylation procedure of Example 252, 82 mg of the amine of Formula U-7, wherein R<sub>1</sub> is ethyl, is reacted with 1-methylimidazole-4-sulfonyl chloride. Flash chromatography on silica using 3% methanol in dichloromethane provides 101 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.8, 1.3, 1.6–2.2, 2.5, 3.5, 3.6, 3.9, 6.8–7.4 ppm HRMS: 555.2192

R<sub>f</sub> 0.29 (5% methanol in dichloromethane)

### **EXAMPLE 255**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula U-8:  $R_1$  is ethyl,  $R_2$  is 5-cyanopyridine-2-yl) Refer to Chart

Using the general sulfonylation procedure of Example 252, 82 mg of the amine of Formula U-7, wherein  $R_1$  is ethyl, is reacted with 5-cyanopyridine-2-sulfonyl chloride. Flash chromatography on silica using 10–15% ethyl acetate in dichloromethane provides 101 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.9, 1.3, 1.6–2.2, 2.5, 3.9, 6.9–7.3, 8.0, 8.1, 8.9 ppm

HRMS: 557.2059

 $R_f$  0.44 (20% ethyl acetate in dichloromethane)

### PREPARATION 87

1,5-Bis-(4-fluorophenyl)-penta-1,4-dien-3-one (Formula V-2) Refer to Chart V

To a rapidly stirred, ambient temperature solution of 10 g of sodium hydroxide in 100 ml of water and 80 ml of ethanol 5 is added a mixture of 12.4 g of 4-fluorobenzaldehyde of Formula V-1 and 2.9 g of acetone. After 45 minutes, the resulting precipitate is filtered off, washed well with water, and dried under vacuum. Recrystallization from ethyl acetate-hexane yields 10.7 g of the title compound as light 10 yellow platelets.

Physical characteristics are as follows: <sup>1</sup>H NMR 86.9–7.2, 7.6–7.7 ppm IR 1653, 1587, 1508, 984, 835 cm<sup>-1</sup> MS: M+ 270 Anal. Found: C, 75.40; H, 4.41 R, 0.35 (dichloromethane) M.P. 152°–154°

#### PREPARATION 88

1,5-Bis-(4-fluorophenyl)-pentane-3-one (Formula V-3) <sup>20</sup> Refer to Chart V

To a solution of 5.41 g of dienone of Preparation 87 (Formula V-2) in 10 ml of THF and 50 ml of methanol is added 2.0 g of magnesium chips. A water bath is used to maintain the temperature of the reaction near ambient. After the magnesium has been consumed, the reaction mixture is partitioned between dichloromethane and dilute HCl, with two additional dichloromethane extractions of the aqueous phase. The combined organic phase is dried over magnesium sulfate and concentrated under reduced pressure. Flash chromatography of the residue on silica using 50% dichloromethane in hexane affords 3.66 g of the title compound as a yellow oil.

Physical characteristics are as follows: <sup>1</sup>H NMR 82.67, 2.85, 6.9, 7.1 ppm IR 2932, 1716, 1603, 1511, 1223, 1159, 828 cm<sup>-1</sup> MS: M+ 274 R<sub>r</sub> 0.28 (50% dichloromethane in hexane)

### PREPARATION 89

4-Hydroxy-5,6-dihydro-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-2-one (Formula V-4) Refer to Chart V

Using the general acetoacetate condensation and ring closure procedure of Preparation 84 (Formula U-4), 3.9 g of the ketone of Preparation 88 (Formula V-3) is converted to 2.86 g of the title compound, which may be recrystallized from dichloromethane-hexane.

Physical characteristics are as follows: <sup>1</sup>H NMR 82.1, 2.57, 2.7, 7.0, 7.1 ppm IR 2924, 1659, 1578, 1508, 1241, 1216 cm<sup>-1</sup> MS: M+ 358 Anal. Found: C, 70.17; H, 5.50 M.P. 140°-141°

#### PREPARATION 90

 $3-[1-(3-Benzyloxycarbonylaminophenyl)-2,2-dimethylpropyl]-6,6-bis[2-(4-fluorophenyl)ethyl]-4-hydroxy-5,6-dihydro-2H-pyran-2-one (Formula V-6: <math>R_1$  is tert-butyl) Refer to Chart V

Using the general benzylidene condensation and cuprate addition procedure of Preparation 85 (Formula U-6), 1.075 g of the dihydropyrone of Preparation 89 (Formula V-4) is converted to 707 mg of the title compound (via the intermediate compound of Formula V-5), which is purified by flash chromatography on silica gel using 40% ethyl acetate in hexane.

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Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 1.07, 2.0, 2.6, 3.9, 5.16, 6.8–7.5 ppm HRMS: 654.3023  $R_f$  0.25 (40% ethyl acetate in hexane)

#### PREPARATION 91

3-[1-(3-Aminophenyl)-2,2-dimethylpropyl]-6,6-bis[2-(4-fluorophenyl)ethyl]-4-hydroxy-5,6-dihydro-2H-pyran-2-one (Formula V-7:  $R_1$  is tert-butyl) Refer to Chart V

Using the general transfer hydrogenolysis procedure of Preparation 86 (Formula U-7), 684 mg of the carbamate of Preparation 90 (Formula V-6, R<sub>1</sub> is tert-butyl) is converted to 497 mg of the title compound, which is purified by flash chromatography on silica gel using 5-10% ethyl acetate in dichloromethane.

Physical characteristics are as follows: 
<sup>1</sup>H NMR δ1.09, 2.0, 2.6, 6.8–7.1 ppm
R, 0.34 (10% ethyl acetate in dichloromethane)
The compound of Formula V-7 wherein R, is

The compound of Formula V-7 wherein  $R_1$  is ethyl is prepared from V-4 by analogous procedures as in the preparation of V-7 wherein  $R_1$  is tert-butyl

(Preparations 90 and 91).

# **EXAMPLE 256**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula V-8:  $R_1$  is tert-butyl,  $R_2$  is 1-methylimidazole-4-yl) Refer to Chart V

Using the general sulfonylation procedure of Example 252, 78 mg of the amine of Preparation 91 (Formula V-7, R<sub>1</sub> is tert-butyl) is reacted with 1-methylimidazole-4-sulfonyl chloride. Flash chromatography on silica using 3-4% methanol in dichloromethane provides 92 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.94, 1.7-2.1, 2.5, 3.50, 6.8-7.4 ppm HRMS: 664.2647 R<sub>f</sub> 0.34 (5% methanol in dichloromethane)

#### **EXAMPLE 257**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula V-8:  $R_1$  is tert-butyl,  $R_2$  is 5-cyanopyridine-2-yl) Refer to Chart V

Using the general sulfonylation procedure of Example 252, 78 mg of the amine of Preparation 91 (Formula V-7, R<sub>1</sub> is tert-butyl) is reacted with 5-cyanopyridine-2-sulfonyl chloride. Flash chromatography on silica using 10–15% ethyl acetate in dichloromethane provides 91.5 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 0.92, 1.9, 2.6, 3.2, 6.8–7.5, 8.0, 8.9 ppm HRMS: 686.2488  $R_{r}$  0.28 (10% ethyl acetate in dichloromethane)

### EXAMPLE 258

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4-fluorophenyl)ethyl)-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula V-8:  $R_1$  is ethyl,  $R_2$  is 1-methylimidazole-4-yl) Refer to Chart V

Using the general sulfonylation procedure of Example 252, 74 mg of the amine of Formula V-7, wherein  $R_1$  is ethyl, is reacted with 1-methylimidazole-4-sulfonyl chloride. Flash chromatography on silica using 3–4% methanol in dichloromethane provides 77 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.87, 2.0, 2.6, 3.62, 4.0, 4.05, 6.9–7.5 ppm HRMS: 636.2350

R<sub>f</sub> 0.31 (5% methanol in dichloromethane)

#### **EXAMPLE 259**

fluorophenyl)ethyl)-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide (Formula V-8: R<sub>1</sub> is ethyl, R<sub>2</sub> is 5-cyanopyridine-2-yl) Refer to Chart V

Using the general sulfonylation procedure of Example 252, 74 mg of the amine of Formula V-7, wherein R, is ethyl, is reacted with 5-cyanopyridine-2-sulfonyl chloride. Flash chromatography on silica using 10% ethyl acetate in dichloromethane provides 83 mg of the title compound as an 15 amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.83, 2.0, 2.6, 3.96, 6.8–7.2, 8.0, 8.8 ppm HRMS: 658.2200

R<sub>c</sub> 0.49 (10% ethyl acetate in dichloromethane)

## **EXAMPLE 260**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2Hpyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula D-6: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) Refer to Chart D

Using the general sulfonylation procedure of Example 252, 54 mg of the amine of Formula D-5 (R<sub>1</sub> and R<sub>2</sub> are propyl, R<sub>3</sub> is tert-butyl), prepared by procedures analogous to those described for the preparation of D-5 (where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) in Preparation 20, is coupled with 5-cyanopyridine-2-sulfonyl chloride of Formula D-7 (R<sub>4</sub> is 5-cyanopyridine-2-yl) to yield, after flash chromatography on silica gel using 10-15% ethyl acetate in dichloromethane, 62 mg of the title compound as an amorphous solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.90, 1.2–1.8, 2.5, 7.0–7.4, 8.1, 8.2, 8.9 ppm HRMS: 525.2305

R<sub>f</sub> 0.44 (20% ethyl acetate in dichloromethane)

## **EXAMPLE 261**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2Hpyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1Himidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 1-methylimidazole-4-yl) Refer 45 to Chart D

Using the general sulfonylation procedure of Example 252, 54 mg of the amine of Formula D-5 (R<sub>1</sub> and R<sub>2</sub> are propyl, R<sub>3</sub> is tert-butyl), is coupled with 1-methylimidazole-4-sulfonyl chloride of Formula D-7 ( $R_4$  is  $^{50}$ 1-methylimidazole-4-yl) to yield, after flash chromatography on silica gel using 3-5% methanol in dichloromethane, 53 mg of the title compound as an amorphous solid.

Physical characteristics are as follows:

MS: 503.2422

R<sub>f</sub> 0.26 (5% methanol in dichloromethane)

#### **EXAMPLE 262**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2Hpyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula D-6: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) Refer to Chart D

Using the general sulfonylation procedure of Example 252, 67 mg of the amine of Formula D-5 (R<sub>1</sub> and R<sub>2</sub> are 65 propyl, R<sub>3</sub> is ethyl) of Preparation 20 is coupled with 5-cyanopyridine-2-sulfonyl chloride of Formula D-7 (R4 is

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5-cyanopyridine-2-yl) to yield, after flash chromatography on silica gel using 10% ethyl acetate in dichloromethane, 78 mg of the title compound as an amorphous solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.6–1.0, 1.2–1.8, 3.4, 3.5, 6.9–7.4, 8.0–8.2, 8.9

HRMS: 498.2072

R, 0.38 (15% ethyl acetate in dichloromethane)

## **EXAMPLE 263**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-phenethyl-6propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) Refer to Chart D

Using the general sulfonylation procedure of Example 252, 79 mg of the amine of Formula D-5 (R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl) of Preparation 20 is coupled with 5-cyanopyridine-2-sulfonyl chloride of Formula D-7 (R4 is 5-cyanopyridine-2-yl) to yield, after flash chromatography on silica gel using 10% ethyl acetate in dichloromethane, 102 mg of the title compound as an amorphous solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–1.0, 1.2–2.6, 3.4, 3.5, 6.9–7.3, 7.9–8.2, 8.9

HRMS: 560.2231

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R<sub>c</sub> 0.37 (15% ethyl acetate in dichloromethane)

#### **EXAMPLES 264-265**

The following compounds are prepared using the general sulfonylation procedure of Example 246. The requisite amine is prepared analogously from the compound of Formula Q-1 (Preparation 69) following Preparations 80 and

264) N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.0, 0.4, 0.6, 1.0, 1.2, 1.7, 2.5, 3.7, 4.1, 6.9–7.6 HRMS: 556.2833 (FAB)

265) N-[3-(1-[6,6-Bis-(2-cyclopropyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran- 3-yl]-2,2-dimethylpropyl)phenyl]-5-cyano-2-pyridinesulfonamide

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.0, 0.4, 0.6, 1.0, 1.2, 1.7, 2.5, 4.1, 7.0–7.5, 8.0, 8.1, 9.0

HRMS: 578.2689 (FAB)

## PREPARATION 92

(3(2E),4S)-3-(2-pentenyl)-4-phenyl-2-oxazolidinone (Formula W-4) Refer to Chart W

A 1 L round-bottomed flask with nitrogen inlet and addition funnel is charged with 6.92 g of (S)-(+)-4-phenyl-2-oxazolidinone and 250 mL of tetrahydrofuran and then cooled to  $-78^{\circ}$  C. To the aforementioned solution is added 25.6 mL of n-butyl lithium during which time a white solid <sup>1</sup>H NMR δ0.9, 0.96, 1.2-1.8, 2.5, 3.6, 3.7, 6.9-7.5 ppm 55 separated from the reaction solution, W-3. To that suspension is added 4.88 g of trans-2-pentencyl chloride of formula W-2 (prepared from the treatment of commercially available trans-2-pentenoyl acid of formula W-1 with oxalyl chloride) in a small volume of THF. The resulting pale yellow homogeneous solution is allowed to warm to room temperature and stirred for another 20 min. The reaction mixture is quenched by the addition of saturated ammonium chloride solution and is extracted with ethyl acetate. The organic layer is separated, washed with brine and water, dried over magnesium sulfate, filtered and concentrated to give a white solid. Recrystallization from hot hexane gives 9.13 g of the title compound.

Physical characteristics are as follows:

MP 86°-88° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.42–7.23, 7.18–7.09, 5.49, 4.70, 4.28, 2.28, 1.08 ppm.

 $[\alpha]_D$ (CHCl<sub>3</sub>)=+109

Anal. found: C, 68.59; H, 6.25; N, 5.70

## PREPARATION 93

(3(3R),4S)-3-[3-(3-Aminophenyl)-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula W-5) Refer to Chart W

A 1 L three-necked, round-bottomed flask with nitrogen inlet and addition funnel is charged with 8.90 g of copper(I) bromide-dimethyl sulfide complex and 125 mL of THF and then cooled to -40° C. To that suspension is added 43 mL of a 1M solution (in THF) of 3-[bis(trimethylsilyl)amino] 15 phenylmagnesium chloride dropwise over 15 minutes. The reaction mixture is warmed 0° C. for 30 minutes and then a 25 mL THF solution containing 8.85 g of (3(2E),4S)-3-(2pentenoyl)-4-phenyl-2-oxazolidinone of Preparation 92 (formula W-4) is added. The reaction mixture is stirred for 20 30 minutes at 0° C. and quenched by the addition of 1N HCl and then the pH readjusted with 1N NaOH to pH 8. The reaction is washed with water, brine and the organic is dried (Na<sub>2</sub>SO<sub>4</sub>). The organic solvent is evaporated in vacuo and the resulting oil chromatographed over 600 g of silica gel, 25 eluting with ethyl acetate/hexane to afford 7.91 g of the title product.

Physical characteristics are as follows:

MP 94°-95° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.28-7.25, 7.07-6.99, 6.60-6.51, <sub>30</sub> 5.38, 4.63, 4.16, 3.52-3.44, 3.10-2.92, 1.65-1.53, 0.76 ppm. IR (mineral oil) 3437, 3355, 1773, 1696, 1605, 1337, 1322, 1299, 1263, 1212, 1096, 1070, 791, 762, 704 cm<sup>-1</sup>.

Anal. found: C, 71.00; H, 6.67; N, 8.17

EI-MS: [M+]=338.

 $[\alpha]_D$  (19.87 mg/2 mL CHCl<sub>3</sub>)=+60°

## PREPARATION 94

3-[3-(3-[Bis(phenylmethyl)amino]phenyl)-1-oxopentyl]-4-phenyl-2-oxazolidinone, (3R)(4S) (Formula W-6) Refer to Chart W

To a mixture of 25 mL of Na<sub>2</sub>CO<sub>3</sub> and 80 mL of methylene chloride was added 7.90 g of (3(3R),4S)-3-[3-(3-aminophenyl)-1-oxopentyl]-4-phenyl-2-oxazolidinone of Preparation 93 (formula W-5) followed by 15.94 g of benzyl bromide. That mixture is heated at 65° C. for 18 bours, the methylene chloride layer separated, dried (Na<sub>2</sub>SO<sub>4</sub>) and solvent evaporated to yield the crude product as a dark viscous oil. That oil is chromatographed over 700 g of silica gel eluding with 25% ethyl acetate/hexane to yield 8.55 g of the title compound.

Physical characteristics are as follows:

MP 92°-3° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.24, 7.02, 6.53, 5.34, 4.59, 4.14, 3.44, 3.07, 2.89, 1.50, 0.64 ppm

Anal. found: C, 78.47; H, 6.68; N, 5.26  $[\alpha]_D$  (19.602 mg/2 mL CHCl<sub>3</sub>)=+32°

# PREPARATION 95

(3R)(4S) 3-[3-(3-[bis(phenylmethyl)amino]phenyl)-2-(2-methyl-1,3-dioxolan-2-yl)-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula W-8) Refer to Chart W

To 25 mL of methylene chloride is added 2.1 g of the amide of formula W-6 of Preparation 94 and the resulting solution cooled to  $-78^{\circ}$  C. under an atmosphere of nitrogen. To that solution is added 872  $\mu$ L of neat TiCl<sub>4</sub> followed by 65 the addition of 732  $\mu$ L of diisopropylethylamine. The resulting mixture is warmed to 0° C. for 30 minutes and then

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cooled back to  $-78^{\circ}$  C. and 1.3 g of 2-methoxy-2-methyl-1,3-dioxolane of formula W-7 and the resulting reaction is warmed to  $0^{\circ}$  C. and stirred for 1 hour, then quenched with saturated ammonium chloride and extracted with methylene chloride. The organic extract is dried (Na<sub>2</sub>SO<sub>4</sub>) and solvent removed in vacuo to afford the crude material. Silica gel chromatography using 100 g of support and eluding with 10% hexane/methylene chloride afforded 1.76 g of the title product.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.36, 7.08, 5.99, 5.42, 4.80, 4.68, 4.60, 4.25, 3.68, 3.57, 3.48, 3.07, 2.90, 1.5, 0.86, 0.54 ppm Anal. found: C, 75.34; H, 6.99; N, 4.87 [α]<sub>D</sub> (18.086 mg/2 mL CHCl<sub>3</sub>)=+25°

#### PREPARATION 96

(3R)(4S)-3-[2-acetyl-3-[3-(bis(phenylmethyl)amino) phenyl]-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula W-9) Refer to Chart W

To 25 mL of tetrahydrofuran and 10 mL of 30% HClO<sub>4</sub> is added 5.0 g of the title compound of Preparation 95 (formula W-8) and the resulting solution stirred at 40° C. for 3 hours. The reaction is neutralized with saturated NaHCO<sub>3</sub> to pH 8 and then extracted with 400 mL of ether. The ether layer is washed with water, brine and then dried (Na<sub>2</sub>SO<sub>4</sub>) and solvent evaporated in vacuo to afford an oil. Chromatography over 300 g of silica gel eluting with 15% acetone/hexane afforded 4.12 g of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.31, 7.08, 6.59, 6.55, 5.42, 4.67, 4.61, 4.22, 3.09, 1.63, 1.56, 0.61 ppm
Anal. found: C, 77.11; H, 6.76; N, 4.98
[α]<sub>D</sub> (20.172 mg/2 mL CHCl<sub>3</sub>)=-10°

## PREPARATION 97

35 (3R)(4S) 3-[2-[1-(3-[bis(phenylmethyl)amino]phenyl) propyl]-5-hydroxy-1,3-dioxo-5-propyloctyl]-4-phenyl-2oxazolidinone (Formula W-10) Refer to Chart W

To 25 mL of methylene chloride is added 1.32 g of the compound of Preparation 96 (formula W-9) and the resulting solution cooled to -78° C. under an atmosphere of nitrogen. To that solution is added 279 µL of TiCl<sub>4</sub> and 450 µL of diisopropylethylamine and stirring continued for 1 hour. To this solution is added 689 µL of heptanone and the reaction temperature raised to 0° C. for 1.5 hours. The reaction is then quenched by the addition of a saturated ammonium chloride solution and the mixture extracted with methylene chloride. The organic extract is washed with saturated NaHCO<sub>3</sub>, dried (Na<sub>2</sub>SO<sub>4</sub>) and evaporated in vacuo to yield the crude product. Chromatography over 100 g of silica gel eluting with 5% hexane/methylene chloride affords 1.16 g of the title compound as an off white foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 7.36, 7.07, 6.58, 6.54, 5.44, 5.24, 4.69, 4.61, 4.27, 3.21, 3.01, 2.48, 1.90, 1.54, 1.15, 0.81, 0.76, 0.58 ppm

Anal. found: C, 76.62; H, 7.63; N, 4.17 [ $\alpha$ ]<sub>D</sub> (15.380 mg/2 mL CHCl<sub>3</sub>)=+16°

## PREPARATION 98

(3S)-3-[1-(3-(Bis(phenylmethyl)amino)phenyl)propyl]-6,6-dipropyl-5,6-dihydro-4-hydroxy-2H-pyran-2-one (Formula W-11) Refer to Chart W

To 10 mL of dry tetrahydrofuran is added 770 mg of the title compound of Preparation 97 (formula W-10) and the resulting solution cooled to 0° C. under an atmosphere of nitrogen. To that solution is added 150 mg of a 60% oil dispersion of sodium hydride and the reaction is warmed to

20° C. and stirring continued for 16 hours. The reaction is quenched with saturated ammonium chloride and extracted with ethyl acetate. The extract is dried and evaporated in vacuo to yield the crude product. Chromatography over 100 g of silica gel eluting with 15% EtOAc/hexane affords 560 mg of the title product.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 7.34, 6.69, 5.87, 4.69, 4.60, 4.09, 2.28, 2.17, 1.89, 1.73, 1.55, 1.32, 0.88 ppm

[Anal. found: C, 79.71; H, 8.07; N, 2.61]  $[\alpha]_D$  (15.998 mg/2 mL CHCl<sub>3</sub>)=-56°]

#### PREPARATION 99

(3S)-3-[1-(3-aminophenyl)propyl]-6,6-dipropyl-5,6-dipydro-4-hydroxy-2H-pyran-2-one (Formula W-12) Refer to Chart W

The title compound of Preparation 98 (formula W-11) ((3R)-3-[1-(3-bis-benzylaminophenyl)propyl]-6,6-bispropyl-5,6-dihydro-4-hydroxypyran-2-one) 110 mg is added to 20 mL of ethyl acetate. To that solution is added 50 mg of 10% Pd/C and the resulting mixture is hydrogenated at 50 psi for 6 hours. The reaction is filtered through celite to yield 83 mg of the title product.

Physical characteristics are as follows:

IR 2957, 2922, 2855, 2871, 2854, 1378, 1605, 1459, <sub>25</sub> 1617, 1262, 1319, 1251, 1282, 1107 cm<sup>-1</sup>.

 $[\alpha]_D$  (6.526 mg/2 mL CH<sub>3</sub>OH)=-34°

#### PREPARATION 100

(4R)3-(1-oxo-2-pentenyl)-4-phenyl-2-oxazolidinone (Formula X-4) Refer to Chart X

A 2-L, three-necked, round-bottomed flask with nitrogen inlet and addition funnel is charged with (R)-(-)-4-phenyl-2-oxazolidinone (31.2 g) and tetrahydrofuran (1.2 L) and cooled to -78° C. The addition funnel is charged with n-butyllithium (1.6M in hexanes, 117 mL), which is added dropwise to the reaction mixture over 20 min. A white precipitate is formed which is X-3. The reaction mixture is stirred for an additional 30 min at -78° C. The addition funnel is then charged with trans-2-pentenoyl chloride of formula X-2, prepared from the acid of formula X-1, (24.4 g) and tetrahydrofuran (50 mL), and this solution is added to the reaction mixture dropwise over 10 min. The resulting pale yellow homogeneous solution is allowed to warm to room temperature and is stirred for another 30 min. The reaction mixture is quenched by the addition of saturated ammonium chloride solution and is extracted with ethyl acetate (2500 mL). The organic layer is separated, washed with brine and water, dried over magnesium sulfate, filtered and concentrated to give 48 g of a white solid. The solid is recrystallized from ethyl acetate (100 mL) and hexane (200  $\,^{50}$ mL) to give 38.0 g the title product as a white solid.

Physical characteristics are as follows:

MP 86°-88° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.42–7.23, 7.18–7.09, 5.49, 4.70, 4.28, 2.28, 1.08 ppm.

IR (mineral oil) 1785, 1764, 1686, 1638, 1349, 1336, 1329, 1257, 1234, 1214, 1087, 1076, 756, 716, 699 cm<sup>-1</sup> EI-MS: [M+]=245.

### PREPARATION 101

(3(3S),4R)-3-[3-(3-aminophenyl)-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula X-5) Refer to Chart X

A 2-L, three-necked, round-bottomed flask with nitrogen inlet and addition funnel is charged with copper(I) bromide-dimethyl sulfide complex (25.1 g) and tetrahydrofuran (250 65 mL) and cooled to -40° C. The addition funnel is charged with 3-[bis(trimethylsilyl)amino]phenylmagnesium chlo-

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ride (1.0M in THF, 122 mL), which is added dropwise to the reaction mixture over 20 min. The reaction mixture is then allowed to warm from -40° C. to -20° C. over 20 min. The addition funnel is charged with 25 g of the title compound of Preparation 100 (formula X-4) and tetrahydrofuran (100 mL), and this solution is added to the reaction mixture dropwise over 30 min at 0° C. The reaction mixture is then stirred for 15 min at 0° C. and quenched by the addition of saturated ammonium chloride solution (adjusted to pH 8 by 10 addition of ammonium hydroxide). The reaction mixture is poured into ether (2 L) and washed with the ammonium chloride solution until the aqueous layer is no longer blue in color. The organic layer is separated, washed with water, dried over magnesium sulfate, filtered and concentrated to give 58 g of a yellow oil. The crude reaction mixture is then stirred at room temperature in a slurry of silica gel (75 g) and methylene chloride (100 mL) for 1 h. The mixture is filtered, washed with methanol, and concentrated to give 49 g of an oil. Column chromatography on 300 g silica (eluting with 10-75% ethyl acetate-hexane, 100% ethyl acetate) yields 30.9 g of a yellow oil. The oil is crystallized from ethyl acetate (75 mL) and hexane (150 mL) to give 21.4 g of the title compound as a white solid.

Physical characteristics are as follows:

MP 94°-97° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.28-7.25, 7.07-6.99, 6.60-6.51, 5.38, 4.63, 4.16, 3.52-3.44, 3.10-2.92, 1.65-1.53, 0.76 ppm. IR (mineral oil) 3437, 3355, 1773, 1696, 1605, 1337, 1322, 1299, 1263, 1212,1096, 1070, 791, 762, 704 cm<sup>-1</sup>. EI-MS: [M+]=338.

#### PREPARATION 102

(3(3S),4R)-3-[3-(3-(phenylmethyl)amino)phenyl)-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula X-6) Refer to Chart X

To a mixture of 80 mL of Na<sub>2</sub>CO<sub>3</sub> and 280 mL of methylene chloride is added 21.0 g of (3(3S),4R)-3-[3-(3-aminophenyl)-1-oxopentyl]-4-phenyl-2-oxazolidinone (formula X-5) of Preparation 101 followed by 23.4 g of benzyl bromide. That mixture is heated at 65° C. for 18 hours, the methylene chloride layer separated, dried (Na<sub>2</sub>SO<sub>4</sub>) and solvent evaporated to yield the crude product as a dark viscous oil. The oil is chromatographed over 700 g of silica gel eluding with 25% ethyl acetate/hexane to yield 31.42 g of the title compound.

Physical characteristics are as follows:

MP 91.8-93.5

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.32, 7.08, 6.60, 5.34, 4.67, 4.15, 3.43, 3.02, 2.91, 1.56, 0.65 ppm

## PREPARATION 103

(3S)(4S)-3-[3-[3-(Bis(phenylmethyl)amino]phenyl]-2-(2-methyl-1,3-dioxolan-2-yl]-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula X-8) Refer to Chart X

To 12 mL of methylene chloride, under nitrogen, is added 1.55 grams of (3(3S),4R)-3-[3-(3-bisbenzylaminophenyl) pentanoyl]-4-phenyl-2-oxazolidinone (formula X-6) of Preparation 102 and the resulting solution cooled to -78° C. To the aforementioned solution is added 646 µl of TiCl<sub>4</sub> followed by the addition of 525 µl of diisopropylethylamine. After stirring at 0° C. for 30 minutes the reaction is cooled back to -78° C. and 886 µl of 2-methoxy-2-methyl-1,3-dioxolane (formula X-7) (also W-7) is added. The reaction is stirred for 1 hour and then quenched by the addition of saturated NH<sub>4</sub>Cl, then saturated NaHCO<sub>3</sub> (pH 8) and finally extraction of the aqueous with both methylene chloride and ethyl ether. Evaporation of solvent affords a viscous oil which is chromatographed over 150 g of silica gel eluting

with 7% hexane/methylene chloride to afford 1.14 g of the title compound.

Physical characteristics are as follows:

IR (mineral oil) 2920, 2954, 2854, 2870, 1776, 1376, 1453, 1196, 699 cm<sup>-1</sup>.

Anal. found: C, 75.27; H, 6.68; N, 4.55

## PREPARATION 104

(3S)(4R) 3-[2-Acetyl-3-[3-[bis(phenylmethyl)amino] phenyl]-1-oxopentyl]-4-phenyl-2-oxazolidinone (Formula 10 1465, 1386, 1363, 1328, 1249, 1260, 696 cm<sup>-1</sup>. X-9) Refer to Chart X

To 15 mL of THF is added 960 mg of (3(3S),4R)-3-[2-(2-methyl-1,3-dioxan-2-yl)-3-(3-bisbenzylaminophenyl) pentanoyl]-4-phenyl-2-oxazolidinone (formula X-8) of Preparation 103. To that solution is then added 4 mL of 30% 15 perchloric acid and the resulting mixture stirred at 40° C. for 2 hours. The reaction is cooled to room temperature and quenced with the addition of excess saturated NaHCO3. The reaction is extracted with 200 mL of ethyl ether, dried (Na<sub>2</sub>SO<sub>4</sub>) and solvent removed in vacuo to yield 981 mg of the crude product. Chromatography over 100 g of silica gel eluding with 10% pentane/methylene chloride affords 854 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.40, 7.08, 6.61, 6.56, 5.41, 4.96, <sub>25</sub> 4.66, 4.61, 4.21, 3.09, 1.63, 1.65, 0.61

IR (mineral oil) 1778, 1718, 1600, 1695, 1452, 1335, 1385, 1200 cm<sup>-1</sup>.

EI-MS: [M+]=560.

Anal. found: C, 76.81; H, 6.59; N, 4.84.

#### PREPARATION 105

(3S)(4R) 3-[2-[1-[3-[bis(phenylmethyl)amino]phenyl]propyl]-5-hydroxy-1,3-dioxo-5-propyloctyl]-4-phenyl-2oxazolidinone (Formula X-10) Refer to Chart X

To 8 mL of methylene chloride under nitrogen is added 35 440 mg of (3(3S),4R)- 3-[2-(acetyl)-3-(3bisbenzylaminophenyl)pentanoyl]-4-phenyl-2oxazolidinone (formula X-9) of Preparation 104 and that solution is cooled to  $-78^{\circ}$  C. To that solution is added 90  $\mu$ l of TiCl<sub>4</sub> followed by the addition of 143  $\mu$ l of diisopropylethylamine. That solution is warmed to 0° C. for 40 minutes and then cooled back to  $-78^{\circ}$  C. at which time 126  $\mu$ l of 4-heptanone is added and the reaction temperature is elevated to 0° C. and stirring continued for 1.5 hours. The reaction is quenced with the addition of saturated NH<sub>4</sub>Cl followed by the addition of saturated NaHCO<sub>3</sub>. The reaction is extracted with methlene chloride (3x60 mL), dried (Na<sub>2</sub>SO<sub>4</sub>) and evaporated in vacuo to yield the crude product as an oil. That material is chromatographed over silica gel (100 g) eluting with 10% pentane/methylene chloride to  $^{50}$ afford 293 mg of the title compound.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.28, 7.07, 6.56, 5.44, 5.24, 4.68, 4.61, 4.26, 3.21, 3.10, 2.48, 1.90, 1.55, 1.21, 0.81, 0.74, 0.58 IR (mineral oil) 2959, 2931, 1779, 1720, 1690, 1600, 55 1494, 1452, 1385, 1359, 1334, 1238, 698 cm<sup>-1</sup>.

## PREPARATION 106

(3R) 3-[1-[3-[bis(phenylmethyl)amino]phenyl]propyl]-5,6dihydro-4-hydroxy-6,6-dipropyl-2H-pyran-2-one (Formula 60 X-11) Refer to Chart X

To 3 mL of THF was added 28 mg of NaH under nitrogen. To that suspension is added 418 mg of (3(3S),4R)-3-[2-((3hydroxy-3-propyl)hexanoyl)-3-(3-bisbenzylaminophenyl) pentanoyl]-4-phenyl-2-oxazolidinone (Formula X-10) of 65 Preparation 105 also in 3 mL of THF at 20° C. The reaction is stirred for 16 hours, cooled to 0° C. and quenched by

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addition of 1N HCl. The reaction is then made basic with the addition of saturated NaHCO<sub>3</sub>. The aqueous is extracted several times with ethyl acetate, the organic extracts dried (Na<sub>2</sub>SO<sub>4</sub>) and solvent is removed in vacuo to yield 518 mg of crude product. Chromatography over silica gel eluting with 15% EtOAc/hexane affords 128 mg of the title compound.

Physical characteristics are as follows:

IR (mineral oil) 2959, 2931, 2873, 1636, 1599, 1451,

EI-MS: [M+]=511.

#### PREPARATION 107

(3R) 3-[1-[3-[amino]phenyl]propyl]-5,6-dihydro-4hydroxy-6,6-dipropyl-2H-pyran-2-one (Formula X-12) Refer to Chart X

The dihydropyrone of formula X-11 ((3R)-3-[1-(3bisbenzylaminophenyl)-propyl]-6,6-bispropyl-5,6-dihydro-4-hydroxypyran-2-one) of Preparation 106, 110 mg, is added to 20 mL of ethyl acetate. To that solution is added 50 mg of 10% Pd/C and the resulting mixture is hydrogenated at 50 psi for 6 hours. The reaction is filtered through celite to yield 83 mg of the title product.

Physical characteristics are as follows:

IR (mineral oil) 2961, 2932, 2873, 1682, 1623, 1604, 1458, 1384, 1369, 1319, 1282, 1259, 1150, 1108 cm<sup>-1</sup>.

EI-MS: [M+]=331

## PREPARATION 108

2-Phenethyl-2-propen-1-ol (Formula BB-2) Refer to Chart

To a cooled (-10° C.) solution of N,N,N,N,-tetramethyl-1,2-ethylenediamine (24.1 mL) in hexane (50 mL) is slowly added butyl lithium (100 mL of a 1.6M solution in hexane). After stirring for 45 minutes at -10° C, the mixture is cooled (-78° C.) and 2-methyl-2-propen-1-ol (BB-1, 6.41 mL) is added dropwise. The reaction is allowed to warm to room temperature and stirred an additional 72 h. The mixture is cooled to -78° C. and a solution of benzyl bromide (8.6 mL) in anhydrous THF (10 mL) is added slowly. The mixture is stirred at -78° C. for 1 hour then gradually allowed to warm to room temperature. After stirring an additional 2 hours, the reaction is quenched by the addition of saturated aqueous NH<sub>4</sub>Cl. The organic layer is diluted with diethyl ether and washed with brine, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo. Purification by flash chromatography using methylene chloride/ethyl acetate/hexane (1:1:6) as eluent affords the title compound (3.5 g) as an oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.31-7.16, 5.07, 4.93, 4.09, 2.82-2.76, 2.41-2.36 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ148.30, 141.69, 128.24, 125.80, 109.76, 65.90, 34.52, 34.16 ppm.

## PREPARATION 109

(2S)-2-Phenethyloxiranemethanol (Formula BB-8) Refer to Chart BB

To a cooled (-20° C.) slurry of molecular sieves (4 Å, crushed and freshly activated, 150 mg) in methylene chloride (1.5 mL) is added diethyl L-tartrate (22 mg) and titanium(IV) isopropoxide (25 mg). The mixture is stirred for 30 min at -20° C. and tert-butyl hydroperoxide (0.84 mL of a 5-6M solution in nonane) is added. After an additional 25 min at -20° C., a solution of allylic alcohol of formula BB-2 (300 mg) of Preparation 108 in methylene chloride (0.5 mL) is slowly added. The mixture is stirred overnight at -20° C. then warmed to -10° C. After an additional 4 hours the reaction is warmed to 0°-5° C. and quenched with the

addition of water (1 mL). After warming to room temperature, stirring is continued for 1 hour and tartrates hydrolysed by the addition of a 30% aqueous NaOH solution saturated with NaCl (0.1 mL). After 30 minutes, the mixture is filtered through Celite and the aqueous phase extracted with several portions of methylene chloride. The combined organic layers are dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo to provide a residue which is purified by flash chromatography using hexane and a gradient of ethyl acetate (10-20%) as eluent to afford the title product of formula BB-3 (223 mg) as an oil. The enantiomeric excess of the reaction is determined to be 86% by analysis of the <sup>1</sup>H NMR (C<sub>6</sub>D<sub>6</sub>) of the Mosher ester formed by the reaction of BB-3 with (S)-(+)-α-methoxy-α-(trifluoromethyl)phenylacetyl chloride (J. A. Dale, D. L. Dull, H. S. Mosher, J. Org. Chem. (1969) 34:2543).

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.34–7.16, 3.83–3.61, 2.89–2.87, 2.72–2.64, 2.17–2.04, 1.89–1.79 ppm

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ141.17, 137.58, 128.49, 128.21, 126.11, 63.00 59.57, 49.92, 33.58, 30.82 ppm.

#### PREPARATION 110

(2S)-2-Phenethyl-2-phenylmethoxymethyloxirane (Formula BB-9) Refer to Chart BB

To a cooled (0°-5° C.) slurry of sodium hydride (124 mg of a 60% suspension in mineral oil) in THF (10 mL) is added 25 alcohol of formula BB-3 (460 mg) of Preparation 109. The mixture is stirred at 0°-5° C. for 5 minutes, allowed to warm to room temperature and stirred an additional 30 minutes. Benzyl bromide (441 mg) is added and the mixture stirred at room temperature overnight. The mixture is quenched with 30 brine (10 mL) and diluted with ethyl ether. The organic layer is washed with brine, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo to provide a residue which is purified by flash chromatography using hexane and a gradient of ethyl acetate (2-5%) as eluent to afford the title product (510 mg) as an 35 oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.34–7.13, 4.59–4.49, 3.64–3.45,

2.75–2.59, 2.19–2.09, 1.94–1.84 <sup>13</sup>C NMR (CDCl<sub>3</sub>) δ141.42, 137.93, 128.58, 128.38, 127.88, 125.94, 73.23, 71.98, 58.21, 50.37, 33.65, 30.83

## PREPARATION 110a

(3S)-1-Phenyl-3-(phenylmethoxymethyl) hexan-3-ol (Formula BB-10) Refer to Chart BB

To a cooled (-45° C.) solution of Li<sub>2</sub>CuCl<sub>4</sub> (0.28 mL of a 0.1M solution in THF) in THF (2 mL) is added ethylmagnesium bromide (0.203 mL of a 3M solution in ethyl ether). The brown solution is stirred at -45° C. for 45 minutes and the epoxide of formula BB-4 (150 mg) of Preparation 110 is added dropwise over ca. 10 minutes. After one hour the reaction is quenched by the addition of saturated aqueous NH<sub>4</sub>Cl and the aqueous layer extracted with ethyl acetate. The combined organic layers are washed with brine, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo to provide a residue which is purified by flash chromatography using hexane/ethyl acetate (5%) as eluent to afford the title product (150 mg) as an oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.38–7.14, 4.54, 3.37, 2.66–2.58, 60

2.20, 1.88–1.76, 1.58–1.52, 1.39–1.25, 0.92

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ142.61, 138.10, 128.41, 128.33, 127.71, 127.63, 125.68, 75.45, 73.79, 73.44, 38.95, 38.52, 29.86, 16.79, 14.37 ppm.

### PREPARATION 110b

(2R)-2-Phenethyl-2-(p-toluenesulfonyloxymethyl) oxirane (Formula BB-13) Refer to Chart BB

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To a cooled (ca. -10° C.) solution of the compound of formula BB-8 (245 mg) of Preparation 109 in methylene chloride (4 mL) is added 4-toluenesulfonyl chloride (302 mg), triethylamine (160 mg) and 4-dimethylaminopyridine (8 mg). The mixture is stirred at ca. -10° C. overnight then warmed to 0°-5° C. for 1 hour. The mixture is diluted with methylene chloride, washed with saturated aqueous NaHCO<sub>3</sub> and brine, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo to provide a residue which is purified by flash chromatography using hexane/ ethyl acetate (5%) as eluent to afford the title compound (448 mg).

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.79, 7.33, 7.28–7.08, 4.13–3.98, 2.64-2.58, 2.44, 2.11-2.00, 1.92-1.82

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ145.17, 140.67, 132.53, 129.96, 128.48, 128.19, 127.95, 126.15, 71.98, 56.49, 50.63, 32.89, 30.37, 21.64 ppm.

#### PREPARATION 110c

(2S)-2-Phenethyl-2-propyl oxirane (Formula BB-12) Refer to Chart BB 20

To a cooled (-45° C.) solution of Li<sub>2</sub>CuCl<sub>4</sub> (0.3 mL of a 0.1M solution in THF) in THF (2 mL) is added ethylmagnesium bromide (0.22 mL of a 3M solution in ethyl ether). The brown solution is stirred at -45° C. for 45 minutes, cooled to -65° C. then to sylate of formula BB-13 (200 mg) of Preparation 110b is added dropwise over ca. 10 minutes. The mixture is stirred for 2.5 hours, warmed to -50° C. for 2 hours and then quenched by the addition of saturated aqueous NH<sub>4</sub>Cl. The aqueous layer is extracted with ethyl acetate and the combined organic layers washed with brine, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo to provide a residue which is purified by flash chromatography using hexane/ethyl acetate (5%) as eluent to afford the title product (60 mg) and hydroxytosylate of formula BB-14 (47

Hydroxytosylate of formula BB-14 is converted to the epoxide of formula BB-12 as follows: To a cooled (0°-5° C.) solution of the compound of formula BB-14 (43 mg) in methanol (2 mL) is added anhydrous K<sub>2</sub>CO<sub>3</sub> (20 mg). After 1 hour at 0°-5° C. the mixture is warmed to room temperature, stirred an additional 90 minutes then quenched by the addition of saturated aqueous NH<sub>4</sub>Cl. The aqueous layer is extracted with ethyl acetate and the combined organic layers washed with brine, dried (MgSO<sub>4</sub>), filtered and concentrated in vacuo to provide a residue which is purified by flash chromatography using hexane/ethyl acetate (5%) to afford the epoxide of formula BB-12 (20 mg).

Physical characteristics for BB-12 are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.31–7.16, 2.68, 2.59, 1.98–1.82, 1.73-1.37, 0.94

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ141.71, 128.41, 128.24, 125.92, 59.11, 52.57, 36.42, 36.02, 31.03, 18.19, 14.22 ppm.

Physical characteristics for BB-14 are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.79, 7.34, 7.29-7.11, 3.90, 2.58-2.53, 2.44, 1.87, 1.77-1.72, 1.54-1.48, 1.31-1.21, 0.89 <sup>13</sup>C NMR (CDCl<sub>3</sub>) δ145.10, 141.66, 132.50, 129.97, 128.45, 128.25, 127.97, 125.96, 74.33, 73.07, 38.37, 37.87, 29.37, 21.66, 16.47, 14.48 ppm.

#### PREPARATION 111

(4S)-3-acetyl-4-phenyl-2-oxazolidinone (Formula FF-3) Refer to Chart FF

To a solution of (S)-(+)-4-phenyl-2-oxazolidinone of formula FF-2 (20 g) in anhydrous tetrahydrofuran (600 mL), cooled to -78° C. is added a solution of 1.6M n-butyllithium in hexanes (77.8 mL) and the resulting suspension stirred at ~78° C. for 30 minutes. The suspension is treated with acetyl

chloride of formula FF-1 (10.23 mL) and then gradually allowed to warm to room temperature. The reaction mixture is quenched with 1 L of saturated ammonium chloride and then partitioned between water and ethyl acetate. The organic layer is separated and the aqueous layer reextracted twice with ethyl acetate. The combined organic layers are washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo. The crude solid is recrystallized from ethyl acetate/hexane affording (21.27 g,) as a white solid.

Physical Characteristics are as follows:

Mp 86°-87° C.

 $^{1}H$  NMR (CDCl<sub>3</sub>)  $\delta 7.42-7.26$ , 5.44-5.40, 4.68, 4.30-4.26, 2.52 ppm

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ169.50, 153.71, 138.81, 128.97, 128.53, 125.73, 69.73, 57.20, 23.59 ppm

#### PREPARATION 112

(3(2E),4S)-3-[4,4-dimethyl (2-pentenoyl)]-4-phenyl-2oxazolidinone (Formula FF-4) Refer to Chart FF

To a solution of the compound of formula FF-3 of Preparation 111 (21.27 g) in anhydrous methylene chloride (500 mL), cooled to -78° C., is added titanium tetrachloride (12.0 mL) in a dropwise manner. The suspension is treated with diisopropylethylamine (19.9 mL) and is allowed to stir at -78° C. for 30 minutes. The suspension is then treated with trimethylacetaldehyde (11.4 mL) followed by diisopropylethylamine (19.9 mL) and allowed to gradually warm to room temperature. After 1 hour the reaction mixture is quenched with water (200 mL) and stirred vigorously for 15 minutes. The organic layer is separated and the aqueous layer is reextracted with methylene chloride. The combined organic layers are washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo. The crude solid is recrystallized from ethyl acetate/hexane affording 21.6 grams of the title compound as a off-white solid:

Physical characteristics are as follows:

Mp 148°-149° C

 $^{1}\text{H}$  NMR (CDCl<sub>3</sub>)  $\delta 7.42-7.05$  , 5.51-5.46, 4.69, 4.30-4.25, 1.09 ppm

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ165.17, 161.61, 153.70, 139.16, 129.12, 128.61, 125.97, 115.71, 69.88, 57.74, 34.31, 28.56

## PREPARATION 113

(3(3S),4S)-3-[3-(3-Aminophenyl)-4,4-dimethylpentanoyl]-4-phenyl-2-oxazolidinone (Formula FF-5) Refer to Chart FF 45

To a slurry of copper(I) bromide dimethylsulfide complex (18.76 g) in anhydrous tetrahydrofuran (60 mL), cooled to -78° C., is added a 1.0M solution of 3-[bis(trimethylsilyl) amino)phenylmagnesium chloride in tetrahydrofuran (182.2) mL) and the resulting slurry stirred at -78° C. for 5 minutes. 50 The slurry is allowed to warm to -15° C. for 15 minutes and then cooled to -78° C. The slurry is then treated with the compound of formula FF-4 of Preparation 112 (16.6 g) added via a solid addition funnel and allowed to stir at -78° C. for 3 hours. The reaction mixture is poured into saturated 55 ammonium chloride (200 mL) and then partitioned between water and ethyl acetate. The organic layer is separated and the aqueous layer (pH 8) is basified to pH 9.5 with concentrated ammonium hydroxide. The aqueous layer is reextracted three times with ethyl acetate, the combined organic 60 layers are washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo. The crude residue is slurried in chloroform (400 mL) and 200 g of silica gel (230-400 mesh) at room temperature for 2 hours. The slurry is filtered and the solids washed several times with chloroform followed by methanol. The filtrate is concentrated in vacuo. Purification by flash chromatography eluting with

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hexane/ethyl acetate (15-40%) afford 17.52 grams of the title compound as a light yellow solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.26-7.12, 7.01, 6.74-6.70, 6.61-6.50, 5.32-5.28, 4.56, 4.11-3.95, 3.48, 2.97-2.91, 0.91

ppm <sup>13</sup>C NMR (CDCl<sub>3</sub>) δ172.34, 153.51, 145.37, 142.24, 153.51 110.80 116.49. 138.12, 128.67, 128.23, 127.68, 124.71, 119.80, 116.49, 113.02, 69.39, 57.39, 52.30, 34.75, 33.49, 27.83 ppm

#### PREPARATION 114

(3(3S),4S)-3-[3-(3-Bisbenzylaminophenyl)-4,4dimethylpentanoyl]-4-phenyl-2-oxazolidinone (Formula FF-6) Refer to Chart FF-6

To a solution of the compound of formula FF-5 of Preparation 113 (15.0 g) in methylene chloride (190 mL) at room temperature is added saturated sodium carbonate (48.7 mL) followed by benzyl bromide (14.3 mL) and the resulting mixture is refluxed for 24 hours. The reaction mixture is allowed to cool to room temperature and partitioned between water (300 mL) and methylene chloride. The organic layer is separated, washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo. Purification by flash chromatography eluting with hexane/ethyl 25 acetate (10-25%) affords 15.1 grams of the title compound as a white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.29–6.99, 6.69–6.49, 5.32–5.26,

4.71-4.50, 4.06-3.94, 2.90-2.81, 0.73 ppm

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ172.79, 153.78, 148.32, 142.08, 139.04, 138.48, 129.07, 128.59, 128.37, 127.89, 126.83, 124.92, 118.44, 114.64, 110.87, 69.70, 57.68, 54.77, 52.97, 34.84, 33.79,27.97 ppm

#### PREPARATION 115

 $^{35}$  [S,[R\*,S\*(E)]]-N-(2-hydroxy-1-methyl-2-phenylethyl)methyl-N-pentenamide (Formula NNN-3) See Chart NNN

A 250-mL, three-necked, round-bottomed flask with nitrogen inlet and addition funnel is charged with the compound of formula NNN-1 (6.6 g) (prepared from the treatment of commercially available trans-2-pentenoic acid with oxalyl chloride) and tetrahydrofuran (80 mL). The addition funnel is charged with a solution of (1R,2S)-ephedrine of formula NNN-2 (7.2 g) and triethylamine (6.0 mL) in tetrahydrofuran (15 mL), which is added dropwise to the reaction mixture. After stirring an additional hour, the reaction mixture is poured into 200 mL of ethyl acetate, washed with three 25-mL portions of water, and concentrated in vacuo to yield 13.5 g of an oil. Column chromatography on 100 g silica (elution with 10-100% ethyl acetate-hexane) yields 10.75 g of the title compound as a colorless oil.

Physical characteristics are as follows:

HRMS found: 248.1652.

## PREPARATION 116

 $[1R-[1R*(R*)2S*]]-3-Amino-\beta-ethyl-N-(2-hydroxy-1$ methyl-2-phenylethyl)-N-methyl-benzenepropanamide (Formula NNN-4) See Chart NNN

A 50-mL, three-necked round-bottomed flask with a nitrogen inlet is charged with the title compound of Preparation 115 (0.247 g) and 5 mL of t-butyl methyl ether and cooled to 0° C. Propyl magnesium chloride (0.55 mL of 2.0M solution in ether) is added dropwise, and the reaction mixture is stirred for an additional 15 min. 3-[Bis(trimethylsilyl) amino phenylmagnesium chloride (2.0 mL of 1.0M solution in tetrahydrofuran, 2.0 mmol) is added dropwise, and the resulting mixture is stirred for an additional 2 h at 0° C. and 1 h at room temperature. The reaction mixture is then

quenched with saturated aqueous ammonium chloride solution (pH adjusted to 8 with ammonium hydroxide) and partitioned between 100 mL of ethyl acetate and 5 mL of water. The organic layer is separated, washed with additional ammonium chloride solution and water, and concentrated in vacuo to give 0.72 g of a yellow oil. The crude oil is then dissolved in chloroform, and silica gel is added to the solution. The resulting mixture is stirred at room temperature for 1.5 h, then filtered through Celite, rinsing with methanol, and concentrated in vacuo to give 0.38 g of a 10 silica gel to give 600 mg of the title compound as an oil. yellow oil. Column chromatography on 50 g of silica gel (elution with 20-100% ethyl acetate-hexane) yields 0.174 g of the title compound as an oil.

Physical characteristics are as follows: HRMS found: 340.2162.

#### PREPARATION 117

[1R-[1R\*(R\*)2S\*]]-3-[bis(phenylmethyl)amino]-b-ethyl-N-(2-hydroxy-1-methyl-2-phenylethyl)-N-methylbenzenepropanamide (Formula NNN-5) See Chart NNN

A 50-mL, three-necked, round-bottomed flask with a condenser fitted with a nitrogen inlet is charged with the title product of Preparation 116 (0.548 g) in 8 mL of acetonitrile. Sodium carbonate (0.375 g) and benzyl bromide (0.42 mL) are added, and the reaction mixture is heated to reflux for 4 25 h. The reaction mixture is then concentrated in vacuo and partitioned between 100 mL of ethyl acetate and 10 mL of water. The organic layer is separated, washed with another 10 mL of water, and concentrated in vacuo to give 1.0 g of a yellow oil. Column chromatography on 65 g of silica gel (elution with 20-100% ethyl acetate-hexane and 5% methanol-methylene chloride) yields 0.447 g of the title compound as a pale yellow oil.

Physical characteristics are as follows:

HRMS found: 520.3102.

#### PREPARATION 118

1-phenyl-6,6,6-trifluoro-3-hexanol (Formula PPP-2) Refer to Chart PPP

To a stirred solution of 4.0 g of ethyl 4,4,4trifluorobutyrate of formula PPP-1 in 25 mL of tetrahydro- 40 furan at -70° C. 24 mL of DiBAL-H (1M in toluene) is added dropwise and the solution stirred for 90 min. In a separate flask containing 680 mg of magnesium turnings and 5 mL of tetrahydrofuran is added 1-phenyl-2-bromoethane in 20 mL of tetrahydrofuran at a rate to maintain reflux. Heating of the mixture is continued for an additional 1 h, then cooled to room temperature and added via cannula to the DiBAL-H reaction above. The resulting white suspension is stirred 30 min at -70° C. and then allowed to warm to room temperature. The reaction is quenched with saturated aqueous ammonium chloride, diluted with 1N hydrochloric acid to dissolve the precipitated salts and extracted with ethyl acetate. The organic layers are combined, washed with water and brine, dried over anhydrous sodium sulfate, filtered and concentrated in vacuo. The crude product is flash 55 chromatographed on silica gel eluting with 20% ethyl acetate in hexane to give 2.0 g of the title compound as a colorless oil.

Physical characteristics are as follows:

HRMS: 232.1088

IR (neat liquid): 3385, 2950, 1455, 1255, 1140, 700 cm<sup>-1</sup>.

#### PREPARATION 119

1-phenyl-6,6,6-trifluoro-3-hexanone (Formula PPP-3) Refer to Chart PPP.

To a solution of 0.48 mL of oxalyl chloride in 10 mL of dichloromethane at -60° C. is added dropwise 0.81 mL of 130

dimethylsulfoxide. The solution is stirred for 5 min then treated with 860 mg of 1-phenyl-6,6,6-trifluoro-3-hexanol of formula PPP-2 of Preparation 118 in 5 mL of dichloromethane and stirred for 15 min. Triethylamine (1.5 mL) is added, the mixture is allowed to warm to room temperature, diluted with water and the layers separated. The aqueous layer is extracted with dichloromethane, the organic layers combined, dried over anhydrous sodium sulfate, filtered, and concentrated. The resulting oil is flash chromatographed on

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>): δ7.2–7.3, 2.9, 2.7, 2.6, 2.4.

<sup>13</sup>C NMR (CDCl<sub>3</sub>): δ206, 140, 129, 128, 126, 125, 44, 35, 30, 28.

#### PREPARATION 120

5,6-dihydro-4-hydroxy-6-phenethyl-6-(3',3',3'trifluoropropyl)-2H-pyran-2-one (Formula PPP-4) Refer to Chart PPP

A suspension of 350 mg of 50% sodium hydride in 10 mL of tetrahydrofuran at 0° C. is treated dropwise with 0.78 mL of methyl acetoacetate. After stirring 30 min, 4.5 mL of 1.6M n-butyllithium in hexane is added and stirring continued for 15 min. A solution of 840 mg of 1-phenyl-6,6,6trifluoro-3-hexanone in 5 mL of tetrahydrofuran of formula PPP-3 of Preparation 119, is added, stirred at 0° C. for 15 min, then allowed to warm to room temperature and stirred for 1 h. The reaction mixture is quenched with saturated aqueous ammonium chloride and extracted with ethyl acetate. The organic layers are washed with water and brine, concentrated in vacuo, then dissolved in 20 mL of tetrahydrofuran. The solution is diluted with 60 mL of water and treated with 20 mL of 1N sodium hydroxide, stirred for 3 h at room temperature, concentrated in vacuo to remove the tetrahydrofuran, cooled to 5° C., and acidified with concentrated hydrochloric acid. The mixture is extracted with dichloromethane, dried over anhydrous sodium sulfate, filtered and concentrated. The crude material is flash chromatographed on silica gel eluting with 30% ethyl acetate in hexane to give 870 mg of the title compound.

Physical characteristics are as follows:

ANAL: C, 61.14, H, 5.45.

## PREPARATION 121

[3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(3, 3,3 -trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl] phenyl-carbamic acid, phenylmethylester (Formula QQQ-3 where R<sub>1</sub>=t-Bu) Refer to Chart QQQ

The title compound of Preparation 120 (850 mg) in 25 mL of tetrahydrofuran at 0° C. is treated with 750 mg of aluminum trichloride, stirred 15 min, and then 700 mg of 3-benzyloxycarbonylaminobenzaldehyde is added. The mixture is allowed to stir at room temperature for 2 h then treated with 2 g of sodium carbonate monohydrate and 0.1 mL of water, stirred for 30 min, and filtered through celite washing the filter cake with tetrahydrofuran. The filtrate is concentrated in vacuo. The resulting material is dissolved in 25 mL of tetrahydrofuran and 285 mg of cuprous bromidedimethylsulfide complex is added and the mixture stirred for 15 min before adding 11 mL of 1M t-butylmagnesium bromide in tetrahydrofuran dropwise over 15-20 min. The resulting brown mixture is stirred an additional 15 min then quenched at 0° C. with 50 mL of water. The layers are separated and the aqueous layer acidified with concentrated hydrochloric acid to dissolve inorganic salts and then extracted with ethyl acetate. The combined organic layers are washed with water and brine, dried over anhydrous sodium sulfate, filtered, and concentrated in vacuo. Flash

chromatography on silica gel eluting with ethyl acetate in hexane gives 1.19 g of the title compound as a white to buff colored foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>): δ7.1–7.7, 6.7, 6.5, 4.4, 1.8–2.8, 1.16. 5 HRMS: 609.2711.

# PREPARATION 122

Preparative resolution of [3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl-carbamic acid, phenylmethylester (Formula QQQ-3 where  $R_1$ =t-Bu) into 4 isomers, 3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethyl-propyl]phenyl-carbamic acid, phenylmethylester (Formulas QQQ-4-7 where  $R_1$  is t-Bu) Refer to Chart QQQ

The first phase of the resolution is accomplished with a 5.1×25 cm (R,R)Whelk-O 1 column eluted with 15% (V/V) isopropanol in hexane at 99 mL/min. ((R,R)Whelk-O 1 is a registered trademark of Regis Technologies, Inc., Morton Grove, Ill. 60053.) The peaks eluting at approximately 54 and 87 min are, respectively, pure Isomer 3 and Isomer 2 as judged from System A, below. The mixture of unresolved Isomer 1 and Isomer 4 eluted at approximately 64 minutes and is further treated as described below.

In the second phase of the resolution, the mixture from above that elutes near 64 minutes is injected onto a  $2.1\times25$  cm Chiralcel OD column (Chiral Technologies, Inc.) and eluted with 35% isopropanol in hexane (V/V) at 8 mL/min. The peaks that elute near 8.7 and 23.9 minutes are, respectively, Isomer 1 and Isomer 4.

In both phase of the resolution of enantiomers, fractions are pooled after assay with System A and pools are concentrated to dryness on a rotary evaporator at 30 mm and a bath 35 set at 50° maximum.

The four constituent enantiomers are (in order of elution from system A) designated (Peak #1), (Peak #2), (Peak #3) and (Peak #4). System A consists of a 0.46×25 cm Chiralcel OD-H column eluted at 1.0 mL/min with 20% isopropanol in hexane (V/V). (Chiralcel OD-H is a registered trademark of Chiral Technologies, Inc., Exton Pa. 19341.)

## PREPARATION 123

3(R or S)-[1-(3-aminophenyl)-2,2-dimethylpropyl]-4-hydroxy-5,6-dihydro-6-(R or S)-phenethyl-6-(3,3,3-trifluoropropyl)-2H-pyran-2-one (Formula QQQ-8, R<sub>1</sub>=t-Bu) Refer to Chart QQQ

A solution of 210 mg of the compound identified as peak 1 from Preparation 122 in 10 mL of methanol is treated with 400 mg of ammonium formate and 40 mg of 10% palladium on charcoal, stirred 2 h, filtered through celite washing the filter cake with methanol. The filtrate is diluted with ethyl acetate and washed with water and brine, dried over anhydrous sodium sulfate, filtered and concentrated in vacuo to give 160 mg of the title compound as a white foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD): δ6.9–7.3, 6.6, 4.1, 2.6–2.7, 1.9–2.4, 1.0.

TLC (silica gel GF): R<sub>f</sub>=0.24 (40% ethyl acetate in hexane).

### **EXAMPLE 266**

5-cyano-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-13, R<sub>1</sub> is t-Bu [R<sub>2</sub> is [5-cyano-2-pyridinyl]) Refer to Chart QQQ

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A solution of the title product of Preparation 123 (50 mg), pyridine (30 mL), and 5-cyano-pyridine-2-sulfonyl chloride (30 mg) in dichloromethane at 0° C. is stirred for 2 h. The crude reaction mixture is chromatographed on silica gel to give the title compound as a white amorphous solid.

Physical characteristics are as follows:

FAB HRMS: 642.2267.

#### EXAMPLE 267

N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula QQQ-13,  $R_1$  is t-Bu, [ $R_2$  is 1-methyl-4-imidazolyl]) Refer to Chart QQQ

Following the procedure described in EXAMPLE 266 and substituting 5-cyano-2-pyridine sulfonyl chloride with 1-methylimidazole-4-sulfonyl chloride the title compound is prepared.

Physical characteristics are as follows:

HRMS: 619.2298

#### **EXAMPLE 268**

5-amino-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-13,  $R_1$ =t-Bu, [ $R_2$  is 5-amino-2-pyridinyl]) Refer to Chart QQQ

A solution of the title product of Preparation 123 (50 mg), pyridine (30  $\mu$ L), and 5-nitro-pyridine-2-sulfonyl chloride (30 mg) in dichloromethane at 0° C. is stirred for 2 h. The crude reaction mixture is chromatographed on silica gel to give the sulfonamide as a white amorphous solid. The white solid is dissolved in 4 mL of methanol and treated with 25 mg of ammonium formate and 5 mg of 10% palladium on carbon, stirred for 1 h at room temperature, diluted with water and extracted with dichloromethane, dried over anhydrous sodium sulfate, filtered, and concentrated in vacuo to give the title compound as an off-white amorphous solid.

Physical characteristics are as follows:

FAB HRMS: 632.2393.

## PREPARATION 124

3(R or S)-[1-(3-aminophenyl)-2,2-dimethyl-propyl]-4-hydroxy-5,6-dihydro-6(S or R)-phenethyl-6-(3',3',3'-trifluoropropyl)-2H-pyran-2-one (Formula QQQ-9, [R<sub>1</sub> is t-Bu]) Refer to Chart QQQ

Following the procedure described in Preparation 123 beginning with the compound isolated from peak 2 from Preparation 122 and using starting materials and reagents known and available to one of ordinary skill in organic synthesis the title compound is prepared.

Physical characteristics are as follows: 1H NMR (CD<sub>3</sub>OD):  $\delta$ 6.9-7.3, 6.6, 4.1, 2.6-2.7, 1.9-2.4, 1.0.

TLC (silica gel GF):  $R_f$ =0.24 (40% ethyl acetate in 55 hexane).

## EXAMPLE 269

5-cyano-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-14,  $R_1$  is t-Bu [ $R_2$  is 5-cyano-2-pyridinyl]) Refer to Chart QQQ

A solution of the title product of Preparation 124 (50 mg), pyridine (30  $\mu$ L), and 5-cyanopyridine-2-sulfonyl chloride (30 mg) in dichloromethane at 0° C. is stirred for 2 h. The crude reaction mixture is chromatographed on silica gel to give the title compound as a white amorphous solid.

Physical characteristics are as follows: FAB HRMS: 642.2260.

## **EXAMPLE 270**

N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4sulfonamide (Formula QQQ-14, R<sub>1</sub> is t-Bu [R<sub>2</sub> is 1-methyl-4-imidazolyl]) Refer to Chart QQQ

Following the procedure described in Example 266 substituting 5-cyano-2-pyridine sulfonyl chloride with 1-methylimidazole-4-sulfonyl chloride the title compound is prepared.

Physical characteristics are as follows:

HRMS: 619.2362

### **EXAMPLE 271**

5-amino-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide 20 (Formula QQQ-14, R<sub>1</sub> is t-Bu [R<sub>2</sub> is 5-amino-2-pyridinyl]) Refer to Chart QQQ

Following the procedure described in Example 268 the title compound is prepared.

Physical characteristics are as follows:

FAB HRMS: 632.2387

## PREPARATION 125

3-(S or R)-[1-(3-aminophenyl)-2,2-dimethyl-propyl]-4hydroxy-5,6-dihydro-6-(R or S)-phenethyl-6-(R or S)-(3',3', 3'-trifluoropropyl)-2H-pyran-2-one (Formula QQQ-10 [R, is t-Bu]) Refer to Chart QQQ

Following the procedure described in Preparation 123 beginning with the compound isolated from peak 3 from known and available to one of ordinary skill in organic synthesis the title compound is prepared.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD): δ6.9–7.3, 6.6, 4.1, 2.6–2.7, 1.9–2.4,

TLC (silica gel GF): R<sub>f</sub>=0.24 (40% ethyl acetate in hexane).

## **EXAMPLE 272**

5-cyano-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-15, R<sub>1</sub> is t-Bu, [R<sub>2</sub> is 5-cyano-2-pyridinyl]) Refer to Chart QQQ

A solution of the title product of Preparation 125 (50 mg), pyridine (30 mL), and 5-cyanopyridine-2-sulfonyl chloride (30 mg) in dichloromethane at 0° C. is stirred for 2 h. The crude reaction mixture is chromatographed on silica gel to give the title compound as a white amorphous solid.

Physical characteristics are as follows:

FAB HRMS: 642.2254

## **EXAMPLE 273**

N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4sulfonamide (Formula QQQ-15, R<sub>1</sub> is t-Bu, [R<sub>2</sub> is 1-methyl-4-imidazolyl]) Refer to Chart QQQ

Following the procedure described in Example 266 substituting 5-cyano-2-pyridine sulfonyl chloride with 1-methylimidazole-4-sulfonyl chloride the title compound is prepared.

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Physical characteristics are as follows: FAB HRMS: 642.2397

**EXAMPLE 274** 

5-amino-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-15, R<sub>1</sub> is t-Bu [R<sub>2</sub> is 5-amino-2-pyridinyl]) Refer to Chart QQQ

Following the procedure described in Example 268 the

10 title compound is prepared.

Physical characteristics are as follows:

FAB HRMS: 632.2393

#### PREPARATION 126

3-(S or R)-[1-(3-aminophenyl)-2,2-dimethyl-propyl]-4-15 hydroxy-5,6-dihydro-6 (S or R)-phenethyl-6-(3',3',3'trifluoropropyl)-2H-pyran-2-one (Formula QQQ-11 [R<sub>1</sub> is t-Bu]) Refer to Chart QQQ

Following the procedure described in Preparation 123 beginning with the compound isolated from peak 4 from Preparation 122 and using starting materials and reagents known and available to one of ordinary skill in organic synthesis the title compound is prepared.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD): δ6.9–7.3, 6.6, 4.1, 2.6–2.7, 1.9–2.4,

TLC (silica gel GF): R=0.24 (40% ethyl acetate in hexane)

#### **EXAMPLE 275**

5-cyano-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-16, R<sub>1</sub> is t-Bu, [R<sub>2</sub> is 5-cyano-2-pyridinyl]) Refer to Chart QQQ

A solution of the title product of Preparation 126 (50 mg), Preparation 122 and using starting materials and reagents 35 pyridine (30 mL), and 5-cyanopyridine-2-sulfonyl chloride (30 mg) in dichloromethane at 0° C. is stirred for 2 h. The crude reaction mixture is chromatographed on silica gel to give the title compound as a white amorphous solid.

Physical characteristics are as follows:

FAB HRMS: 642.2248.

## **EXAMPLE 276**

N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4sulfonamide (Formula QQQ-16, R<sub>1</sub> is t-Bu, [R<sub>2</sub> is 1-methyl-4-imidazolyl]) Refer to Chart QQQ

Following the procedure described in Example 266 substituting 5-cyano-2-pyridine sulfonyl chloride with 1-methylimidazole-4-sulfonyl chloride the title compound is prepared.

Physical characteristics are as follows:

FAB HRMS: 620.2403

## **EXAMPLE 277**

5-amino-N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide (Formula QQQ-16, R<sub>1</sub> is t-Bu, [R<sub>2</sub> is 5-amino-2-pyridinyl]) Refer to Chart QQQ

Following the procedure described in Example 268 the title compound is prepared.

Physical characteristics are as follows:

FAB HRMS: 632.2406

## **EXAMPLE 278**

65 N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2phenethyl)-[6-(R or S)-propyl]-2H-pyran-3-yl]-2,2dimethylpropyl]phenyl]-2-pyridinesulfonamide

Following procedures analogous to those described above and using Isomer 2 of Preparation 143 the title compound is prepared.

Physical characteristics are as follows: FAB HRMS: 562.2527.

#### **EXAMPLE 279**

N-[3-(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R or S)-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide

Following procedures analogous to those described above and using Isomer 1 of Preparation 143 the title compound is prepared.

Physical characteristics are as follows:

#### FAB HRMS: 562.2528.

## PREPARATION 127

2-Mercapto-4-trifluoromethylpyridine

To 1.0 g of 2-chloro-4-trifluoromethylpyridine (Lancaster Chemical Co) is added 10 ml of absolute ethanol and 417 mg of thiourea. The reaction mixture is heated at reflux for 4 hours and 1.25 ml of a solution of 7.44 g KOH in 20 ml of water is added. The solution is heated at reflux for an additional 1 hour. The reaction solution is cooled and poured into 100 ml of a 0.1N NaOH solution. The resulting solution is extracted three times with 100 ml of methylene chloride and the resulting aqueous solution is acidified to pH 4 by addition of glacial acetic acid. The aqueous solution is extracted three times with 100 ml of methylene chloride and the organic solution is dried over anhydrous sodium sulfate. Filtration followed by evaporation to dryness gives 501 mg of a yellow crystalline solid.

Physical characteristics are as follows: Found C:40.22; H:2.33; N:8.07; S:17.59

HRMS: 179.0019

## PREPARATION 128

2-Chlorosulfonyl-4-trifluoromethylpyridine

To 425 mg of 2-mercapto-4-trifluoromethylpyridine of Preparation 127 is added 10 ml of 1N aqueous HCl. The reaction mixture is cooled to 0° C. and Cl<sub>2</sub> gas is bubbled into the cold reaction mixture for 15 minutes. The reaction mixture is filtered and the resulting solid is washed well with water. The white solid is dissolved in methylene chloride and is washed twice with saturated aqueous NaHCO<sub>3</sub> followed by one wash with water. After drying the organic solution over sodium sulfate (anhydrous), the solution is filtered and evaporated to dryness to give 300 mg of 2-chlorosulfonyl-4-trifluoromethylpyridine which is used directly without further purification, and stored at -78° C. until ready for use.

#### PREPARATION 129

2-Chlorosulfonyl-5-trifluoromethylpyridine

Substituting 2-mercapto-5-trifluoromethylpyridine for 2-mercapto-4-trifluoromethylpyridine in the reaction above in Preparation 128 gives 2-chlorosulfonyl-5-trifluoromethylpyridine as a colorless oil which slowly crystallizes. This material is used without further purification and stored at -78° C. until ready for use.

## PREPARATION 130

3-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl]phenyl-carbamic acid, (Formula SSS-1;  $R_1$  is ethyl; Refer to Chart SSS)

To 7.2 g of AlCl<sub>3</sub> at -70° C., under N<sub>2</sub>, is added 180 ml 65 of THF. The mixture is allowed to stir at 0° C. for 15 minutes and 5.38 g of Formula SSS-A; Refer to Chart SSS, prepared

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by procedures analogous to those described in Preparation 17, is added. The reaction mixture is stirred for 15 minutes and 6.88 g of 3-aminoCbZ-benzaldehyde (Formula SSS-B; Refer to Chart SSS) is added. The reaction mixture is stirred for 15 minutes at 0° C. followed by 3 hours at room temperature. The reaction is cooled to 0° C. and 35 g of sodium carbonate monohydrate is added, with vigorous stirring, followed by 1.6 ml of water. After stirring at 0° C. for an additional 15 minutes, 120 ml of THF is added and the mixture filtered through celite. The celite is washed well with THF and the THF solution is evaporated to dryness under vacuum to an amber foam. The residue is dissolved in 180 ml of THF, the solution is cooled to −5° C. and 3.2 g of CuBr.Me<sub>2</sub>S added. The mixture is stirred for 15 minutes and 15 65 ml of a 2M ethylmagnesium chloride in THF solution is added, dropwise, with temperatures not rising above 0° C. The reaction is allowed to stir for an additional 15 minutes and 9 ml of water is slowly added followed by 45 ml of 1N HCl. These additions are done at 0° C. The reaction mixture is poured into 2 L of ethyl ether and 200 ml of water is added. The aqueous layer is separated and the organic layer is extracted three times with 10% aqueous ammonium carbonate followed by once with water. The organic solution is dried over anhydrous sodium sulfate, filtered and evaporated to dryness to give 10.2 g of a crude amorphous foam. This crude material is chromatographed over silica gel using 2% ethyl acetate in methelene chloride as eluent to give 4.74 g of 3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl]phenyl-carbamic acid.

## PREPARATION 130A

 $3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-phenethyl-6-propyl)-2H-pyran-3-yl]-propyl]-phenyl-carbamic acid (Formula RRR-1; <math>R_1$  is ethyl; Refer to Chart RRR)

Following the procedure of Preparation 130 beginning with the compound from Preparation 17 the title compound is prepared.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD): δ6.9–7.5, 5.1, 4.0, 1.4–2.7, 0.9.

TLC (silica gel GF): Rf=0.28, 30% ethyl acetate in hexane.

## PREPARATION 131

Preparative chiral resolution of 3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl] phenyl-carbamic acid (Formula SSS-1; R<sub>1</sub> is ethyl; Refer to Chart SSS) to give two isomers of 3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl]phenyl-carbamic acid (Formulas SSS-3 and SSS-4; R<sub>1</sub> is ethyl; Refer to Chart SSS).

Samples of the title compound of Preparation 130 are injected onto a 2.1×25 cm Chiralcel OD column and eluted with 20% isopropanol (V/V) in hexane at 10 mL/min. The material eluting near 19.1 minutes is 3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl]phenyl-carbamic acid, ( $\alpha$ )25D +26° (methanol), (Formula SSS-3; Refer to Chart SSS) (peak 1) and that eluting near 37.7 minutes is 3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl] phenyl-carbamic acid ( $(\alpha)^{25}_D$  -27° (methanol), (Formula SSS-4; Refer to Chart SSS) (peak 2). The pools are concentrated separately on a rotary evaporator (ca. 30 mm, bath at 50° maximum) to give white amorphous solids.

## PREPARATION 132

3(R or S)-[1-(3-aminophenyl)-propyl]-4-hydroxy-5,6-dihydro-6,6-dipropyl-2H-pyran-2-one (Formula SSS-5; R<sub>1</sub> is ethyl; Refer to Chart SSS)

To 1.04 g of 3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6,6-di-n-propyl-2H-pyran-3-yl]-propyl]phenyl-carbamic acid (Formula SSS-3; Refer to Chart SSS) of Preparation 131, the compound identified as peak 1 from the chiral resolution of the product of Preparation 131, is added 20 ml of methanol and 1.29 g of ammonium formate. When dissolution is complete, 275 mg of 10% Pd/C is added and the reaction mixture is stirred at room temperature for 60 minutes. The reaction mixture is filtered (celite) and the methanolic solution is evaporated to dryness to give a crude 10 solid. The crude solid is partitioned between methylene chloride and water. The organic layer is washed twice with water and dried over anhydrous sodium sulfate. The methylene chloride solution is filtered and evaporated to dryness to give 625 mg of 3(R or S)-[1-(3-aminophenyl)-propyl]-4- 15 hydroxy-5,6-dihydro-6,6-dipropyl-2H-pyran-2-one as an amorphous foam.

Physical characteristics are as follows: MS(EI): 331 ( $\alpha$ )<sup>25</sup><sub>D</sub> +38° (c=0.3715, methanol).

#### PREPARATION 133

3(R or S)-[1-(3-aminophenyl)-propyl]-4-hydroxy-5,6-dihydro-6,6-di-n-propyl-2H-pyran-2-one (Formula SSS-6;  $R_1$  is ethyl; Refer to Chart SSS)

To 825 mg of 3(R or S)-[1-(3-aminophenyl)-propyl]-4-hydroxy-5,6-dihydro-6,6-dipropyl-2H-pyran-2-one (Formula SSS-4; Refer to Chart SSS), of Preparation 131, the compound identified as peak 2 from the chiral resolution of the product of Preparation 131, is added 20 ml of methanol and 1.02 g of ammonium formate. When dissolution is complete, 210 mg of 10% Pd/C is added and the reaction mixture is stirred at room temperature for 60 minutes. The reaction mixture is filtered (celite) and the methanolic solution is evaporated to dryness. The crude solid is partitioned between methylene chloride and water. The organic layer is washed twice with water and dried over anhydrous sodium sulfate. The methylene chloride solution is filtered and evaporated to dryness to give 483 mg of title compound as an amorphous foam.

Physical characteristics are as follows: MS(EI): 331

 $(\alpha)^{25}_{D}$  -39° (c=0.2680, methanol).

## **EXAMPLE 280**

5-Trifluoromethyl-N-[3-[1-[4-hydroxy-2-oxo-6,6-di-n-45 propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula SSS-9;  $R_1$  is ethyl;  $R_2$  is 5-trifluoromethylpyridine; Refer to Chart SSS)

The title compound of Preparation 130 is deprotected as in Preparation 132 to give the compound of formula SSS-2. 50 To 132 mg of formula SSS-2 is added 15 ml of methylene chloride and 66 microliters of pyridine. The reaction solution is cooled to -5° C. and 98 mg of 2-chlorosulfonyl-5trifluoromethylpyridine (product of Preparation 129) is added. After stirring at 0° C. for 60 minutes the solution is 55 placed on a silica gel column and eluted with 10% ethyl acetate in methylene chloride until the 5-trifluoromethyl-N-[3-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2Hpyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide is collected. Rf=0.6 in 10% ethyl acetate in methylene chloride. 60 Evaporation of the organic solution to dryness gives 177 mg of 5-trifluoromethyl-N-[3-[1-[4-hydroxy-2-oxo-6,6-di-npropyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2pyridinesulfonamide

Physical characteristics are as follows: MS(EI): 540, 497, 411, 401, 383, 342, 331, 197, 174, 146, 133.

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HRMS: 540.1938

Rf=0.6 in 10% ethyl acetate in methylene chloride.

<sup>1</sup>H NMR(MeOD): 88.91, 8.21-8.19, 7.12, 6.98-6.96, 6.86-6.83, 3.85-3.79, 2.46, 2.10-1.98, 1.84-1.75, 1.58-1.47, 1.27-1.15, 0.82-0.72 ppm

#### **EXAMPLE 281**

5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula SSS-7;  $R_1$  is ethyl;  $R_2$  is 5-trifluoromethylpyridine; Refer to Chart SSS)

To 66 mg of the title product from Preparation 132 (Formula SSS-5; Chart SSS) is added 8 ml of methylene chloride and 33 microliters of pyridine. The reaction solution is cooled to -5° C. and 49 mg of 2-chlorosulfonyl-5-trifluoromethylpyridine (product of Preparation 129) is added. After stirring at 0° C. for 60 minutes the solution is placed on a silica gel column and eluted with 10% ethyl acetate in methylene chloride until the 5-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-di-ydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide is collected. Rf=0.6 in 10% ethyl acetate in methylene chloride. Evaporation of the organic solution to dryness gives 69 mg of the title compound.

Physical characteristics are as follows:

MS(EI): 540, 497, 411, 401, 383, 342, 331, 197, 174, 146, 133.

Rf=0.6 in 10% ethyl acetate in methylene chloride

<sup>1</sup>H NMR(MeOD): 88.91, 8.21-8.19, 7.12, 6.98-6.96,
6.86-6.83, 3.85-3.79, 2.46, 2.10-1.98, 1.84-1.75,
1.58-1.47, 1.27-1.15, 0.82-0.72 ppm

## **EXAMPLE 282**

5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula SSS-8; R<sub>1</sub> is ethyl; R<sub>2</sub> is 5-trifluoromethylpyridine; Refer to Chart SSS)

Following the procedure of Example 281 but substituting the product of Preparation 133 (formula SSS-6) gives 5-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-40 di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

MS(EI): 540, 497, 411, 401, 383, 342, 331, 197, 174, 146, 33.

Rf=0.6 in 10% ethyl acetate in methylene chloride

<sup>1</sup>H NMR(MeOD): 88.91, 8.21-8.19, 7.12, 6.98-6.96, 6.86-6.83, 3.85-3.79, 2.46, 2.10-1.98, 1.84-1.75, 1.58-1.47, 1.27-1.15, 0.82-0.72 ppm

#### **EXAMPLE 283**

4-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula SSS-7;  $R_1$  is ethyl;  $R_2$  is 4-trifluoromethylpyridine; Refer to Chart SSS)

Following the procedure of Example 281 but substituting the product of Preparation 128 for the pyridylsulfonylchloride gives 4-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

MŠ(EI): 146, 145, 139, 133, 71, 57, 55, 43, 41 HRMS: 540.1902

#### **EXAMPLE 284**

4-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula SSS-8;  $R_1$  is ethyl;  $R_2$  is 4-trifluoromethylpyridine; Refer to Chart SSS)

Following the procedure of Example 282 but substituting the product of Preparation 128 for the pyridylsulfonylchloride gives 4-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows: MS(EI): 146, 145, 139, 133, 71, 57, 55, 43, 41 HRMS: 540.1896

#### **EXAMPLE 285**

5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-6;  $R_1$  is t-butyl;  $R_2$  is n-propyl;  $R_3$  is 5-trifluoromethyl-2-pyridinyl; Refer to Chart TTT)

Following the procedure of Example 281 but using Isomer 1 of Preparation 144 (Formula TTT-4; Chart TTT;  $R_1$  is t-butyl,  $R_2$  is n-propyl) gives 5-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows: MS(EI): 163, 162, 147, 146, 69, 57, 56, 43, 41 HRMS: 568.2213

#### **EXAMPLE 286**

5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]1-2-pyridinesulfonamide (Formula TTT-7;  $R_1$  is t-butyl;  $R_2$  is n-propyl;  $R_3$  is 5-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 281 but substituting Isomer 2 of Preparation 144 (Formula TTT-5; Chart TTT;  $R_1$  is t-butyl;  $R_2$  is n-propyl) gives 5-trifluoromethyl-N-[3-[1(R or S)-4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows: HRMS: 568.2237

#### **EXAMPLE 287**

5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-6;  $R_1$  is ethyl;  $R_2$  is phenyl;  $R_3$  is 5-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 281 but using Isomer 1 of Preparation 145 (Formula TTT-4; Chart TTT; R<sub>1</sub> is ethyl; R<sub>2</sub> is phenyl) gives 5-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows:

MS(EI): 665, 647, 456, 455, 333, 134, 133, 117, 105, 91

# EXAMPLE 288

5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-55 diphenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-7; R<sub>1</sub> is ethyl; R<sub>2</sub> is phenyl; R<sub>3</sub> is 5-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 281 but substituting Isomer 2 of Preparation 145 (Formula TTT-5; Chart TTT; R<sub>1</sub> 60 139 is ethyl; R<sub>2</sub> is phenyl) gives 5-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows:

HRMS: 665.2300

MS(EI): 665, 647, 456, 455, 333, 134, 133, 117, 105, 91

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#### **EXAMPLE 289**

4-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-6;  $R_1$  is ethyl;  $R_2$  is phenyl;  $R_3$  is 4-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 281 but substituting Isomer 1 of Preparation 145 (Formula TTT-4; Chart TTT; R<sub>1</sub> is ethyl; R<sub>2</sub> is phenyl) gives 5-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows:

MS(EI): 666, 665, 647, 134, 133, 117, 105, 91 HRMS: 665.2306

## **EXAMPLE 290**

4-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-7;  $R_1$  is ethyl;  $R_2$  is phenyl;  $R_3$  is 4-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 281 but substituting Isomer 2 of Preparation 145 (Formula TTT-5; Chart TTT; R<sub>1</sub> is ethyl; R<sub>2</sub> is phenyl) gives 4-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows:

HRMS: 665.2306

MS(EI): 666, 665, 647, 134, 133, 117, 105, 91

### **EXAMPLE 291**

4-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-6;  $R_1$  is t-butyl;  $R_2$  is methyl;  $R_3$  is 4-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 283 but substituting Isomer 1 of Preparation 144 (Formula TTT-4; Chart TTT;  $R_1$  is t-butyl;  $R_2$  is methyl) gives 4-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows:

MS(EI): 525, 512, 428, 411, 302, 284, 258, 146, 57

HRMS: 568.2209

## **EXAMPLE 292**

4-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide (Formula TTT-7;  $R_1$  is t-butyl;  $R_2$  is methyl;  $R_3$  is 4-trifluoromethylpyridine; Refer to Chart TTT)

Following the procedure of Example 283 but substituting Isomer 2 of Preparation 144 (Formula TTT-5; Chart TTT;  $R_1$  is t-butyl;  $R_2$  is methyl) gives 4-trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide as an amorphous solid.

Physical characteristics are as follows:

MS(EI): 569, 551, 511, 493, 439, 371, 360, 303, 284, 161, 139

HRMS (MI+H+): 569.2297

## PREPARATION 134

N-[3-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl] phenylcarbamic acid, phenylmethyl ester (Formula RRR-1;  $R_1$  is t-butyl; Refer to Chart RRR)

To 4.8 g of AlCl<sub>3</sub> at  $-70^{\circ}$  C., under N<sub>2</sub>, is added 120 ml of THF. The mixture is allowed to stir at  $0^{\circ}$  C. for 15 minutes and 4.68 g of (formula RRR-A) of Preparation 17 is added. The reaction mixture is stirred for 15 minutes and 4.59 g of 3-aminoCbZ-benzaldehyde (formula RRR-B) is added. The reaction mixture is stirred for 15 minutes at 0° C. followed by 3 hours at room temperature. The reaction is cooled to 0° C. and 26 g of sodium carbonate monohydrate (0.21M) is added, with vigorous stirring, followed by 1.08 ml of water. After stirring at 0° C. for an additional 15 minutes, the mixture is treated with 120 ml of THF and filtered through celite. The celite is washed well with THF and the THF solution is evaporated to dryness under vacuum to an amber foam. The residue is dissolved in 120 ml of THF, the solution is cooled to -5° C. and 2.1 g of CuBr.Me<sub>2</sub>S added. The mixture is stirred for 15 minutes and 65 mL of a 1M t-butylmagnesium chloride in THF solution is added, dropwise, with temperatures not rising above 0° C. The reaction is allowed to stir for an additional 15 minutes at 0° C. and 6 ml of water is slowly added followed by 30 ml of 1N HCl. The reaction mixture is poured into 1.3 L of ethyl ether. The aqueous layer is separated and the organic layer is extracted three times with 10% aqueous ammonium carbonate followed by once with water. The organic solution is dried over anhydrous sodium sulfate, filtered and evaporated to dryness to give an amorphous foam. This crude material is chromatographed over silica gel using 30% ethyl acetate in hexane as eluent to give 6.15 g the title product.

#### PREPARATION 135

Preparative resolution of N-[3-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl-carbamic acid, phenylmethyl ester (Formula RRR-1;  $R_1$  is t-butyl) into four isomers, 3-(R or S)-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(R or S)-(2-phenethyl)-6-(R or S)-n-propyl-2H-pyran-3-yl]-2,2-35 dimethyl-propyl]phenyl-carbamic acid, phenylmethyl ester (Formulas RRR-3 to 6;  $R_1$  is t-butyl; Refer to Chart RRR)

The four constituent enantiomers are (in order of elution from system A) Isosmer 1 (Formula RRR-4; refer to Chart RRR), Isomer 2 (Formula RRR-3; refer to Chart RRR), 40 Isomer 3 (Formula RRR-5; refer to Chart RRR), and Isomer 4 (Formula RRR-6; refer to Chart RRR). System A consists of a 0.46×25 cm Chiralcel OD-H column eluted at 0.5 mL/min with 20% isopropanol and 0.1% trifluoracetic acid in hexane (V/V). (Chiralcel OD-H is a registered trademark 45 of Chiral Technologies, Inc., Exton Pa. 19341.)

The first phase of the resolution is accomplished with a 2.1×25 cm (R,R)Whelk-O 1 column eluted with 20% (V/V) isopropanol in hexane at 12 mL/min. ((R,R)Whelk-O 1 is a registered trademark of Regis Technologies, Inc., Morton Grove, Ill. 60053.) The peaks eluting at approximately 35 and 41 min are, respectively, a mixture of Isomer 3 and Isomer 4 and a mixture of Isomers 1 and 2 and as judged from System A, above. The two mixtures are further treated as below.

In the second phase of the resolution, the mixture from above that elutes near 41 minutes is injected onto a 2.1×25 cm Chiralcel OD column (Chiral Technologies, Inc.) and elutes with 15% isopropanol and 0.05% trifluoracetic acid in hexane (V/V) at 9.0 mL/min. The peaks that elute near 11.0 60 and 22.0 minutes are designated respectively, peaks 1 and 2 and as judged from System A.

In the final phase of the resolution, the mixture that elutes from the Whelk-O column near 35 minutes is injected onto a 2.2×25 cm Chiralcel OD column and elutes with 35% 65 isopropanol and 0.1% trifluoroacetic acid (V/V) in hexane at 9.0 mL/min. The isomer that elutes near 9.7 minutes is

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designated peak 3 and the one that elutes near 16.6 minutes is designated peak 4.

#### PREPARATION 136

3-[1-(3-Aminophenyl)-2,2-dimethyl-propyl]-5,6-dihydro-4hydroxy-6-phenethyl-6-n-propyl-pyran-2-one (Formula RRR-2, Refer to Chart RRR)

To 590 mg of 3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2phenethyl)-6-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl] phenyl-carbamic acid of Preparation 134, is added 10 ml of methanol and 660 mg of ammonium formate. When all the reactants are dissolved, 140 mg of 10% Pd/C is added and the reaction is allowed to stir at room temperature for 60 minutes. The reaction is filtered (celite) and the filter pad is washed well with methanol and the methanol solution is evaporated under vacuum to a crude solid. The solid is partitioned between water and methylene chloride, and the methylene chloride layer is washed twice with water, dried over anhydrous sodium sulfate and evaporated to dryness to give 372 mg of 3-[1-(3-amino-pheny)-2,2-dimethyl-propyl] -5,6-dihydro-4-hydroxy-6-phenethyl-6-n-propyl-pyran-2one. This material is identical to material described earlier (Formula T-4; refer to Chart T).

#### PREPARATION 137

3(R or S)-[1-(3-amino-phenyl)-2,2-dimethyl-propyl]-5,6-dihydro-4-hydroxy-6(R or S)-phenethyl-6(R or S)-propyl-pyran-2-one (Formula RRR-7; R<sub>1</sub> is t-butyl; Refer to Chart RRR)

Following the procedure of Preparation 136 but substituting the compound in Preparation 135 designated peak 2 for the compound of Preparation 134 gives 3(R or S)-[1-(3-amino-pheny)-2,2-dimethyl-propyl]-5,6-dihydro-4-hydroxy-6(R or S)-phenethyl-6(R or S)-propyl-pyran-2-one as an amorphous foam.

Physical characteristics are as follows:

MS(EI): 421, 365, 164, 163, 147, 146, 118, 107, 91, 57. HRMS: 421.2617

## PREPARATION 138

3(R or S)-[1-(3-Amino-pheny)-2,2-dimethyl-propyl]-5,6-dihydro-4-hydroxy-6(R or S)-phenethyl-6(R or S)-propyl-pyran-2-one (Formula RRR-8;  $R_1$  is t-butyl; Refer to Chart RRR)

Following the procedure of Preparation 136 but substituting the compound of Preparation 135 designated peak 1 for the compound of Preparation 134 gives 3(R or S)-[1-(3-amino-pheny)-2,2-dimethyl-propyl]-5,6-dihydro-4-hydroxy-6(R or S)-phenethyl-6(R or S)-propyl-pyran-2-one as an amorphous foam.

Physical characteristics are as follows:

MS(EI): 421, 365, 164, 163, 147, 146, 118, 107, 91, 57.

## **EXAMPLE 293**

5-Trifluoromethyl-N-[3-[1-[5,6-dihydro-4-hydroxy-2-oxo-55 6-(2-phenethyl)-6-n-propyl-2H-pyran-3-yl]-2,2dimethylpropyl]-phenyl]-2-pyridinesulfonamide (Formula RRR-15; R<sub>1</sub> is t-butyl; R<sub>2</sub> is 5-trifluoromethyl; Refer to Chart RRR).

Following the procedure of Example 281 but substituting the product of Preparation 136 for the product of Preparation 132 gives 5-trifluoromethyl-N-[3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

MŚ(EI): 497, 411, 401, 383, 343, 331, 197, 174, 146, 133 HRMS: 540.1938

## **EXAMPLE 294**

5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2Hpyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2pyridinesulfonamide (Formula RRR-11; R<sub>1</sub> is t-butyl; R<sub>2</sub> is 5-trifluoromethyl; Refer to Chart RRR)

Following the procedure of Example 281 but substituting the product of Preparation 137 for the product of Preparation 132 gives 5-trifluoromethyl-N-[3 (R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n- 10 propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

MS(EI): 373, 355, 201, 146, 145, 118, 117, 91, 57.

HRMS:630.2394

#### **EXAMPLE 295**

5-Trifluoromethyl-N-[3 (R or S)-[1-[5,6-dihydro-4hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2pyridinesulfonamide (Formula RRR-12; R<sub>1</sub> is t-butyl; R<sub>2</sub> is 5-trifluoromethyl; Refer to Chart RRR)

Following the procedure of Preparation 281 but substituting the product of Preparation 138 for the product of Preparation 132 gives 5-trifluoromethyl-N-[3(R or S)-(-(1-25 [5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows: MS(EI): 373, 355, 201, 146, 145, 118, 117, 91, 57.

HRMS:630.2379

### **EXAMPLE 296**

4-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2Hpyran-3-y1]-2,2-dimethylpropy1]-pheny1]-2pyridinesulfonamide (Formula RRR-11, R<sub>1</sub> is t-butyl; R<sub>2</sub> is 4-trifluoromethyl; Refer to Chart RRR)

Following the procedure of Example 294 but substituting the product of Preparation 128 for the product of Preparation 129 gives 4-trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-npropyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

HRMS: 631.2444

## **EXAMPLE 297**

4-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-50 pyran-3-y1]-2,2-dimethylpropy1]-pheny1]-2pyridinesulfonamide (Formula RRR-12; R<sub>1</sub> is t-butyl; R<sub>2</sub> is 4-trifluoromethyl; Refer to Chart RRR)

Following the procedure of Example 295 but substituting the product of Preparation 128 for the product of Preparation 55 129 gives 4-trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-npropyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

MS(EI): 633, 632, 631, 614, 613, 346, 201, 146, 91, 57.

HRMS: 631.2450

## **EXAMPLE 298**

5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-65 2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2Hpyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide

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(Formula RRR-11; R<sub>1</sub> is ethyl; R<sub>2</sub> is 5-trifluoromethyl; Refer to Chart RRR)

Following the procedure of Example 294 but substituting the product of Preparation 147A gives 5-trifluoromethyl-N-[3(R or S)-(-(1-5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-6)]phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-propyl) phenyl]-2-pyridinesulfonamide as an amorphous foam.

Physical characteristics are as follows:

MS(EI): 605, 604, 603, 602, 586, 585, 393, 201, 133, 91 HRMS: 603.2153

#### **EXAMPLE 298A**

5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide (Formula RRR-12; R<sub>1</sub> is ethyl; R<sub>2</sub> is 5-trifluoromethyl; Refer to Chart RRR)

Following the procedure of Example 294 but substituting the amine derived from Isomer 1 of Preparation 147 (derived following the procedure of Preparation 147A) gives the title compound as an amorphous foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD): δ8.9, 8.2, 8.0, 7.0–7.3, 3.9, 2.4–2.7, 1.2–2.2, 0.8–1.0.

TLC (silica gel GF): Rf=0.19, 40% ethyl acetate in hexane.

## PREPARATION 139

(5-Nitro-pyridin-2-yl)-isothiourea hydrochloride (Formula UUU-2) Refer to Chart UUU

A solution of 3.81 g of thiourea in 75 mL of hot absolute ethanol is treated with 7.61 g of 2-chloro-5-nitropyridine (Formula UUU-1) and is heated at reflux for 6 hours. The mixture is then cooled to 0° C. and the precipitated solid is collected. The solid is washed sequentially with cold absolute ethanol and chloroform. The solid is dried in vacuo to afford 6.91 g of the title product as a light brown solid.

Physical characteristics are as follows:

MP 175° C. (dec.)

<sup>1</sup>H NMR (CD<sub>3</sub>OD) δ7.9, 8.6, 9.4 ppm

## PREPARATION 140

5-Nitro-2-thiopyridine (Formula UUU-3) Refer to Chart UUU

A solution of 1.65 g of sodium carbonate in 50 mL of MS(EI): 633, 632, 631, 614, 613, 346, 201, 146, 91, 57 45 water is treated with 2.35 g of the title compound of Preparation 139. The mixture is charged with a solution of 2.75 g of sodium hydroxide in 50 mL of water and the resulting mixture is warmed to room temperature. After stirring for 1 hour, the mixture is heated to 95° C. for 1 hour and finally cooled to room temperature. The aqueous mixture is extracted with two portions of diethyl ether and then carefully acidified with 6N aqueous hydrochloric acid. The orange precipitated solid is collected and washed sequentially with cold dilute aqueous hydrochloric acid and water. The solid is dried in vacuo to afford 1.27 g of the title product as an orange solid.

Physical characteristics are as follows:

MP 167°-170° C.

<sup>1</sup>H NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ7.4, 7.9, 8.5 ppm

# PREPARATION 141

5-Nitro-2-pyridinesulfonyl chloride (Formula UUU-4) Refer to Chart UUU

To a suspension of 1.27 g of the title compound of Preparation 140 in 25 mL of 1N aqueous hydrochloric acid and 5 mL of acetic acid at 0° C. is vigorously bubbled in chlorine gas. After 15 minutes, the chlorine gas addition is

ceased and replaced with nitrogen gas. The resulting solid is collected and washed sequentially with cold dilute aqueous hydrochloric acid and water. The solid is dried in vacuo to afford 1.60 g of the title product as a tan solid.

Physical characteristics are as follows: MP 77°-80° C. <sup>1</sup>H NMR (CDCl<sub>3</sub>) 88.3, 8.8, 9.6 ppm

#### PREPARATION 142

N-[3-(1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-5-nitro-2-pyridinesulfonamide (Formula UUU-5:  $R_1$  is 2-phenylethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl) Refer to Chart UUU

To a solution of 210 mg of the title compound of Preparation 81 (Formula T-4) in 2 mL of dichloromethane at 0° C. is added 80  $\mu$ L of pyridine followed by 111 mg of the title compound of Preparation 141 (Formula UUU-4). After warming to room temperature overnight, the reaction mixture is column chromatographed on flash silica gel eluting 20 with 3% to 9% ethyl acetate in dichloromethane to provide 303 mg of the title compound as a yellow foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ0.8–1.0, 1.2–1.4, 1.6–1.9, 2.4–2.7, 4.0, 6.9–7.4, 8.0, 8.5, 9.4 ppm

HRMS 608.2412 (EI)

## **EXAMPLE 299**

5-Amino-N-[3-(1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl) phenyl]-2-pyridinesulfonamide (Formula UUU-6:  $R_1$  is 2-phenylethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl) Refer to Chart UUU

To a solution of 300 mg of the title compound of Preparation 142 (Formula UUU-5) in 5 mL of methanol under argon is added 500 mg of ammonium formate followed by 100 mg of 10% palladium on carbon. After 1 hour, the reaction mixture is filtered through a pad of Celite with methanol washes. The combined filtrates are concentrated under reduced pressure and the residue is repeatedly triturated with portions of dichloromethane. The combined dichloromethane washes are concentrated under reduced pressure and the residue is column chromatographed on flash silica gel eluting with 50% ethyl acetate in dichloromethane to provide 246 mg of the title compound as a 45 white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD) δ0.8–1.0, 1.2–1.4, 1.5–2.0, 2.4–2.6, 4.0, 6.7, 6.8–7.3, 7.4, 7.9 ppm

HRMS 577.2617 (EI)

## **EXAMPLE 300-327**

Following the procedures and preparations described above and using starting materials known and available to one of ordinary skill in organic synthesis, the following <sup>55</sup> additional compounds in Table 3 of the present invention are made from the compounds prepared in the following preparations:

## PREPARATION 143

Preparative separation of N-[3-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl-carbamic acid phenylmethyl ester, to give 4 isomers

The title compound of Preparation 134 is separated into 65 four constituent steroisomers which are (in order of elution from system A) 4 isomers: Isomer 1, Isomer 2, Isomer 3, and

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Isomer 4 with the following approximate observed retention times 10.5, 14.9, 21.4 and 65.2 minutes respectively. System A consists of a 0.46×25 cm Chiralcel OD-H column eluting at 0.5 mL/min with 20% isopropanol and 0.1% trifluoracetic acid in hexane (V/V). (Chiralcel OD-H is a registered trademark of Chiral Technologies, Inc., Exton Pa. 19341.)

The first phase of the separation is accomplished with a 2.1×25 cm (R,R)Whelk-O 1 column eluting with 20% (V/V) isopropanol in hexane at 12 mL/min. ((R,R)Whelk-O 1 is a registered trademark of Regis Technologies, Inc., Morton Grove, Ill. 60053.) The peaks eluting at approximately 35 and 41 min are, respectively, a mixture of Isomers 3 and 4 and a mixture of Isomers 1 and 2 as judged from System A, above. The two mixtures are further treated as below.

In the second phase of the separation, the mixture from above that elutes near 41 minutes is injected onto a 2.1×25 cm Chiralcel OD column (Chiral Technologies, Inc.) and eluting with 15% isopropanol and 0.05% trifluoracetic acid in hexane (V/V) at 9.0 mL/min. The peaks that elute near 11.0 and 22.0 minutes are, respectively, Isomer 1 and Isomer 2 as judged from System A.

In the final phase of the separation, the mixture that elutes from the (R,R)Whelk-O 1 column near 35 minutes is injected onto a 2.2×25 cm Chiralcel OD column eluting with 35% isopropanol and 0.1% trifluoroacetic acid (V/V) in hexane at 9.0 mL/min. The isomer that elutes near 9.7 minutes is Isomer 3 and the one that elutes near 16.6 minutes is Isomer 4.

In both phases of the separation of stereoisomers, fractions are pooled after assay with System A and pools are concentrated to dryness on a rotary evaporator.

## PREPARATION 144

Resolution of N-[3-(1-[5,6-Dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) phenyl]carbamic acid, phenylmethyl ester to give 2 isomers

Samples of the starting compound (up to 1.0 gm each run) are injected onto a 5.1×50 cm Chiralcel OD column (Chiral Technologies, Inc.). The enantiomers elute at about 23 min (This corresponds to the benzyloxycarbonyl protected analogue of amine (Isomer 1) (EI-MS: 359 [M+]; <sup>1</sup>H NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD): 7.1–6.9, 6.5, 4.2, 2.6–2.3, 1.8–1.2, 1.1, 0.9; TLC: R=0.42 (10% ethyl acetate in dichloromethane)), and at about 33 min (This corresponds to the benzyloxycarbonyl protected analogue of amine (Isomer 2) (EI-MS: 359 [M+]; <sup>1</sup>H NMR (CDCl<sub>3</sub>—CD<sub>3</sub>OD): 7.1–6.9, 6.5, 4.2, 2.6-2.3, 1.8-1.2, 1.1, 0.9; TLC: R<sub>f</sub>=0.42 (10% ethyl acetate in dichloromethane)). The mobile phase is 20% isopropanol and 0.1% acetic acid in hexane (V/V) pumped at 60 mL/min. The purity is checked on a 0.46×25 cm Chiralcel OD-H column (Chiral Technologies, Inc.). The mobile phase is 20% isopropanol in hexane (V/V) and 0.05% trifluoroacetic acid pumped at 0.5 mL/min. The observed retention times are 8.9 and 16.7 min (monitor set at 238 nm) for Isomer 1 and Isomer 2, respectively.

## PREPARATION 145

Resolution of N-[3-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]carbamic acid, phenylmethyl ester to give 2 isomers

Samples of the starting compound (up to 1.3 gm each run) are injected onto a 5.1×50 cm Chiralcel OD column (Chiral Technologies, Inc.). The enantiomers are eluted with 20% isopropanol and 0.025% acetic acid in hexane (V/V) at 60 mL/min until the first enantiomer elutes. At this point (approximately 120 min into the run) the flow rate is increased to 90 mL/min to expedite elution of the second enantiomer. The enantiomers elute near 91.2 min (This is the

corresponding benzyloxycarbonyl analogue of amine Isomer 1 and near 132 min (This is the corresponding benzyloxycarbonyl analogue of amine Isomer 2. The purity is checked on a 0.46×25 cm Chiralcel OD-H column. The mobile phase is 30% isopropanol in hexane (V/V) pumped at 0.5 mL/min.

## PREPARATION 146

5-Carbamoylpyridine-2-sulfonyl chloride (Formula VVV-2) Refer to Chart VVV

Into a cold (0°), stirred suspension of 400 mg of 2-mercapto-5-carbamoylpyridine of formula VVV-1 in 7.5 ml of 1N HCl is passed a brisk stream of chlorine gas. After ten minutes, the suspension is filtered, and the solid washed well with water and dried in vacuo. Obtained is 517 mg of 15 ppm. the title compound as a nearly white solid.

#### **EXAMPLE 328**

 $N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-4-4-4))}]$ fluorophenyl)ethyl-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl]phenyl]-4-cyanobenzenesulfonamide (Formula U-8: R<sub>1</sub> is tert-butyl, R<sub>2</sub> is 4-cyanophenyl) Refer to Chart U

Using the general sulfonation procedure of Example 252, 88 mg of the amine of Preparation 86 (Formula U-7, R, is tert-butyl) is reacted with 4-cyanobenzenesulfonyl chloride. Flash chromatography on silica gel using 10% ethyl acetate in dichloromethane provides 117 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.90, 1.3, 1.7, 2.5, 3.6, 6.8–7.4, 7.6, 7.8 ppm HRMS: 605.2478

R<sub>c</sub> 0.36 (10% ethyl acetate in dichloromethane)

#### **EXAMPLE 329**

 $N-[3-\{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-35))\}]$ fluorophenyl)ethyl-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl}phenyl]-8-quinolinesulfonamide (Formula U-8: R<sub>1</sub> is tert-butyl, R<sub>2</sub> is 8-quinolyl) Refer to Chart U

Using the general sulfonylation procedure of Example 252, 88 mg of the amine of Preparation 86 (Formula U-7, R<sub>1</sub> is tert-butyl) is reacted with 8-quinolinesulfonyl chloride. Flash chromatography on silica gel using 5-10% ethyl acetate in dichloromethane provides 101 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.63, 0.9, 1.1, 1.3, 1.6–1.9, 2.4–2.6, 6.7–7.6, 8.0, 8.2, 9.1 ppm

HRMS: 631.2638

R<sub>f</sub> 0.30 (5% ethyl acetate in dichloromethane)

# **EXAMPLE 330**

 $N-[3-\{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2$ phenylethyl)-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1Himidazole-4-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 1-methylimidazole-4-yl) 55 Refer to Chart D

Using the general sulfonylation procedure of Example 252, 77 mg of the amine of Formula D-5, wherein R<sub>1</sub> and R<sub>2</sub> are phenethyl and R<sub>3</sub> is ethyl, is reacted with 1-methylimidazole-4-sulfonyl chloride. Flash chromatogra- 60 phy on silica gel using 3% methanol in dichloromethane provides 97.0 mg of the title compound as a crystalline white

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.88, 1.9–2.2, 2.6, 3.6–3.8, 3.97, 6.9–7.5 ppm 65 HRMS: 600.2521

R<sub>c</sub> 0.31 (5% methanol in dichloromethane)

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**EXAMPLE 331**  $N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-dihyd$ phenylethyl)-2H-pyran-3-yl)propyl}phenyl]-5-

cyanopyridine-2-sulfonamide (Formula D-6: R<sub>1</sub> is phenethyl, R<sub>2</sub> is phenethyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is

5-cyanopyridine-2-yl) Refer to Chart D

Using the general sulfonylation procedure of Example 252, 77 mg of the amine of Formula D-5, wherein R<sub>1</sub> and R<sub>2</sub> phenethyl and R<sub>3</sub> is ethyl, is reacted with 5-cyanopyridine-10 2-sulfonyl chloride. Flash chromatography on silica gel using 10% ethyl acetate in dichloromethane provides 88.3 mg of the title compound as a crystalline white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.85, 1.8–2.2, 2.5–2.7, 3.97, 6.9–7.4, 7.9, 8.8

HRMS: 622.2355

R<sub>f</sub> 0.28 (10% ethyl acetate in dichloromethane)

## **EXAMPLE 332**

 $N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-dihydro-2-oxo-6))}]$ fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-carbamoylpyridine-2-sulfonamide (Formula U-8: R<sub>1</sub> is ethyl, R<sub>2</sub> is 5-carbamoylpyridine-2-yl) Refer to Chart U

Using the general sulfonylation procedure of Example 252, 82 mg of the amine of Formula U-7, wherein R<sub>1</sub> is ethyl, is reacted with 5-carbamoylpyridine-2-sulfonyl chloride of Preparation 146. Flash chromatography on silica gel using 3-6% methanol in dichloromethane provides 55.4 mg of the title compound as an amorphous solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–0.9, 1.3, 1.6–2.1, 2.5, 3.9, 6.8–7.3, 7.8, 8.2

HRMS: 596.2216

R<sub>f</sub> 0.16 (5% methanol in dichloromethane)

## **EXAMPLE 333**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-(4fluorophenyl)ethyl)-2H-pyran-3-yl)propyl}phenyl]-5carbamoylpyridine-2-sulfonamide (Formula V-8: R<sub>1</sub> is ethyl, R<sub>2</sub> is 5-carbamoylpyridine-2-yl) Refer to Chart V

Using the general sulfonylation procedure of Example 252, 98 mg of the amine of formula V-7, wherein  $R_1$  is ethyl) is reacted with 5-carbamoylpyridine-2-sulfonyl chloride of Preparation 146. Flash chromatography on silica using 3-6% methanol in dichloromethane provides 58.3 mg of the title compound as an amorphous solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.83, 1.8–2.2, 2.5–2.6, 6.8–7.2, 7.8, 8.1, 9.0

HRMS: 676,2297

R<sub>f</sub> 0.17 (5% methanol in dichloromethane)

## **EXAMPLE 334**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2Hpyran-3-yl)propyl}phenyl]-5-carbamoylpyridine-2sulfonamide (Formula D-6: R<sub>1</sub> is propyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R4 is 5-carbamoylpyridine-2-yl) Refer to Chart D

Using the general sulfonylation procedure of Example 252, 66 mg of the amine of Formula D-5 (R<sub>1</sub> and R<sub>2</sub> are propyl, R<sub>3</sub> is ethyl) is coupled with 5-carbamoylpyridine-2sulfonyl chloride of Preparation 146 to yield, after flash chromatography on silica gel using 3-6% methanol in dichloromethane, 83.8 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–0.9, 1.2–2.1, 3.87, 7.0–7.3, 7.8, 8.2 ppm. HRMS: 516.2156

R<sub>f</sub> 0.22 (5% methanol in dichloromethane)

Resolution of N-[3-[1-(4-Hydroxy-5,6-dihydro-2-oxo-6phenethyl-6-propyl-2H-pyran-3-yl)propyl]phenyl]carbamic acid, phenylmethyl ester to give 4 isomers (Formula WWW-2: R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is 5 ethyl) Refer to Chart WWW and RRR

The four isomers of the product of Preparation 130A (Formula RRR-1; R<sub>1</sub>=ethyl) are (in order of increasing retention time on System B): (ca. 16.9 min) (Isomer 1), (ca. 28.0 min) (Isomer 2), (ca. 38.2 min) (Isomer 3) and (ca. 49.8 10 of Preparation 147A. The title compound is obtained as an min) (Isomer 4). System B consists of a 0.46×25 cm Chiralcel OD-H column (Chiral Technologies, Inc.) eluting with 25% isopropanol in hexane (V/V) at 0.5 mL/min.

In Phase one of the complete resolution repeatedly inject 55 mg samples of the product of Preparation 130A onto a 15 ppm. 2.1×25 cm (R,R)Whelk-O 1 column (Regis Technologies, Inc.). Elute the isomers at 10 mL/min with 35% isopropanol and 0.5% acetic acid in hexane (V/V). The first of the three peaks to elute (near 12 min) is a mixture of Isomers 1 (Formula RRR-4 of Chart RRR) and 2 (Formula RRR-3 of 20 Chart RRR) as shown by injecting aliquots in System B. Resolve this mixture in Phase 2, below.

The second phase consists of a 2.1×25 cm Chiralcel OD column kept at 30°. Inject 60 mg batches of the mixture obtained in the first phase and elute the enantiomers with 25 25% isopropanol and 0.05% trifluoroacetic acid (V/V) at 9 mL/min. Separately pooled and concentrated, the fractions eluting near 14.5 and 23.9 min to give Isomers 1 (Formula RRR-4 where R<sub>1</sub> is ethyl of Chart RRR) and 2 (formula RRR-3 where  $R_1$  is ethyl of Chart RRR) respectively.

### PREPARATION 147A

3-(R or S)-[1-(3-aminophenyl)-propyl]-4-hydroxy-5,6dihydro-6-(R or S)-phenethyl-6-(R or S)-propyl-2H-pyran-2-one (Formula RRR-7; R<sub>1</sub> is ethyl; Refer to Chart RRR)

Following the procedure of Preparation 132 beginning with the peak identified as peak 2 (Formula RRR-3; R<sub>1</sub> is ethyl of Chart RRR) from the chiral resolution of the product of Preparation 147, the title compound is prepared.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD): δ6.5–7.3, 3.9–4.0, 2.5–2.7, 1.2–2.3,

TLC (silica gel GF): Rf=0.31, 40% ethyl acetate in hexane.

## **EXAMPLE 335**

 $N-[3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } S)$ S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide (Formula WWW-4: R, is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) [Isomer 1] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the first stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral 55 HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:

8.8-9.0 ppm.

HRMS: 560.2210

R<sub>f</sub> 0.41 (15% ethyl acetate in dichloromethane)

### **EXAMPLE 336**

 $N-[3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S \text{ or }$ R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-

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cyanopyridine-2-sulfonamide (Formula WWW-4: R, is phenethyl, R2 is propyl, R3 is ethyl, R4 is 5-cyanopyridine-2-yl) [Isomer 2] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine that is the title product of Preparation 147A (Formula WWW-3, where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl). The amine is derived from the second stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.6–2.6, 3.3–3.6, 6.9–7.3, 7.7–8.2, 8.8–9.0

HRMS: 560.2215

R<sub>f</sub> 0.41 (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 337**

 $N-[3-\{1(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } R)$ S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide (Formula WWW-4: R<sub>1</sub> is phenethyl, R2 is propyl, R3 is ethyl, R4 is 5-cyanopyridine-2-yl) [Isomer 3] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the third stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral 30 HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.6–2.6, 3.3–3.6, 6.9–7.3, 7.7–8.2, 8.8–9.0

HRMS: 560.2210

 $R_f 0.41$  (15% ethyl acetate in dichloromethane)

## **EXAMPLE 338**

 $N-[3-\{1(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-$ R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide (Formula WWW-4: R, is phenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) [Isomer 4] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the fourth stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–1.0, 1.2–2.6, 3.3–3.6, 6.9–7.3, 7.7–8.2, 8.8-9.0 ppm.

HRMS: 560.2210

R<sub>f</sub> 0.41 (15% ethyl acetate in dichloromethane)

## **EXAMPLE 339**

 $N-[3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-$ <sup>1</sup>H NMR δ0.7-1.0, 1.2-2.6, 3.3-3.6, 6.9-7.3, 7.7-8.2, <sup>60</sup> S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1methyl-1H-imidazole-4-sulfonamide (Formula WWW-4: R, is phenethyl, R2 is propyl, R3 is ethyl, R4 is 1-methylimidazol-4-yl) [Isomer 1] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the first stereoisomer

of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows: <sup>1</sup>H NMR  $\delta$ 0.7-2.8, 3.2-3.7, 3.9, 7.0-7.6 ppm. HRMS: 537.2317 R<sub>f</sub> 0.36 (5% methanol in dichloromethane)

#### **EXAMPLE 340**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or 10 R)-phenethyl- 6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 1-methylimidazol-4-yl) [Isomer 2] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is phenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the second stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3-4% methanol in dichloromethane.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 0.7-2.7, 3.3-3.7, 4.0, 7.0-7.5 ppm. HRMS: 537.2275  $R_f$  0.36 (5% methanol in dichloromethane)

#### **EXAMPLE 341**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 1-methylimidazol-4-yl) [Isomer 3] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> is phenethyl, R<sub>2</sub> is propyl, and 35 R<sub>3</sub> is ethyl. The amine is derived from the third stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows: 

<sup>1</sup>H NMR δ0.7–2.7, 3.3–3.7, 4.0, 7.0–7.5 ppm. HRMS: 537.2329

# $R_f$ 0.36 (5% methanol in dichloromethane)

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4:  $R_1$  is phenethyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 1-methylimidazol-4-yl) [Isomer 4] Refer to Chart WWW

**EXAMPLE 342** 

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is phenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the fourth stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 147. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows: <sup>1</sup>H NMR 80.7-2.8, 3.2-3.7, 3.9, 7.0-7.6 ppm. HRMS: 537.2312 R<sub>f</sub> 0.36 (5% methanol in dichloromethane)

## PREPARATION 148

3-[(3-Nitrophenyl)methyl]-6,6-diphenethyl-4-hydroxy-5,6-65 dibydro-2H-pyran-2-one (Formula XXX-3) Refer to Chart XXX

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To a solution of 172 mg of 6,6-Diphenethyl-4-hydroxy-5,6-dihydro-2H-pyran-2-one of formula XXX-1 and 81 mg of meta-nitrobenzaldehyde in 2 ml of dry THF, under argon, is added a solution of 142 mg of AlCl<sub>3</sub> in 1 ml of THF. The solution is stirred at room temperature for 2 hours, then quenched with 310 mg of sodium carbonate decahydrate, diluted with ether, and filtered through Celite with ether rinses. Following removal of solvent under reduced pressure, 264 mg of crude benzylidene of Formula XXX-2 is obtained. This material is dissolved in 5 ml of methanol, and the solution cooled to 0° for the addition of 44 mg of sodium cyanoborohydride. After an hour, a further 20 mg aliquot of sodium cyanoborohydride is added. After another 30 minutes, the mixture is acidified with dilute HCl to pH 1 and extracted with three portions of dichloromethane. The extract is dried (MgSO<sub>4</sub>) and then concentrated under reduced pressure. Flash chromatography of the residue on silica using 5-20% ethyl acetate in dichloromethane provides 211 mg of the title compound as an amorphous white 20 solid.

Physical characteristics are as follows:  $^1H$  NMR  $\delta2.0$ , 2.7, 3.8, 7.0–7.4, 7.6, 8.0, 8.2 ppm. MS: M+ 457 R<sub>f</sub> 0.25 (5% ethyl acetate in dichloromethane)

#### PREPARATION 149

3-[(3-Aminophenyl)methyl]-6,6-diphenethyl-4-hydroxy-5, 6-dihydro-2H-pyran-2-one (Formula XXX-4) Refer to Chart XXX

A mixture of 211 mg of the product of Preparation 148 (Formula XXX-3) and 50 mg of 10% palladium on carbon in 5 ml of methanol is stirred at room temperature under 1 atmosphere hydrogen gas. After two hours, the mixture is filtered through Celite and concentrated under reduced pressure. Flash chromatography of the residue on silica using 25% ethyl acetate in dichloromethane affords 133.6 mg of the title compound.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta$ 2.0, 2.6, 3.6, 4.1, 6.5, 6.6, 6.7, 6.9–7.3 ppm. MS: M+ 427  $R_{f}$  0.33 (25% ethyl acetate in dichloromethane)

## **EXAMPLE 343**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula YYY-5)  $R_1$  and  $R_2$  are phenethyl,  $R_3$  is 1-methylimidazole-4-yl) Refer to Chart YYY

Using the general sulfonylation procedure of Example 252, 77 mg of the amine of formula YYY-4 ( $R_1$  and  $R_2$  are phenethyl) is reacted with 1-methylimidazole-4-sulfonyl chloride. Flash chromatography on silica using 3% methanol in dichloromethane provides 90.7 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.96, 1.0, 1.6–2.7, 3.45, 6.8–7.5 ppm.

HRMS: 628.2832

R<sub>f</sub> 0.38 (3% methanol in dichloromethane)

# **EXAMPLE 344**

60 N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula YYY-5) R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is 5-cyanopyridine-2-yl) Refer to Chart YYY

Using the general sulfonylation procedure of Example 252, 77 mg of the amine of Formula YYY-4 (R<sub>1</sub> and R<sub>2</sub> are phenethyl) is reacted with 5-cyanopyridine-2-sulfonyl chlo-

ride. Flash chromatography on silica using 10% ethyl acetate in dichloromethane provides 86.1 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.96, 1.8–2.2, 2.5–2.8, 4.1, 4.3, 6.9–7.4, 5 7.9–8.0, 8.9 ppm.

HRMS: 650.2681

R<sub>f</sub> 0.27 (10% ethyl acetate in dichloromethane)

## PREPARATION 150

Resolution of N-{3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-(2-(4-fluorophenyl)ethyl-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl]phenyl]carbamic acid, phenylmethyl ester to give 4 isomers (Formula WWW-2: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is t-butyl) Refer to Chart 15

System C is used to track the enantiomers and to monitor the preparative columns. System C consists of a 0.46×25 cm Chiralcel OD-H column (Chiral Technologies, Inc.) with 15% isopropanol in hexane (V/V) at 0.5 mL/min. The peaks eluting near 13.5, 18.8, 37.1 and 79.7 min are, respectively, Isomer 1, Isomer 2, Isomer 3, and Isomer 4.

Separate Isomers 3 and 4 from the mixture on a 2.1×25 cm (R,R)Whelk-O 1 column (Regis Technologies, Inc.). These two isomers elute at about 23.9 and 26.8 min when the column is developed with 20% isopropanol in hexane (V/V) at 10 mL/min at 30°. The desired isomers elute as an unresolved mixture near 28.9 min and are separated in the second stage of the resolution.

For the second stage inject the unresolved mixture onto a  $2.1\times25$  cm Chiralcel OD column (Chiral technologies, Inc.) kept at 30°. With 12% isopropanol in hexane (V/V) at 12 mL/min, Isomer 1 emerges near 14.5 min and Isomer 2 emerges near 20.8 min.

## **EXAMPLE 345**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl,  $R_4$  is 5-cyanopyridine-2-yl) [Isomer 1] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the first stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.7–2.7, 3.2, 3.5, 3.6, 3.7, 4.1, 6.8–7.4, 7.5, 7.8–8.2, 8.8 ppm.

HRMS: 606.2429

R<sub>f</sub> 0.40 (15% ethyl acetate in dichloromethane)

## **EXAMPLE 346**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide 60 (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl,  $R_4$  is 5-cyanopyridine-2-yl) [Isomer 2] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of 65 Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the

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second stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR 80.6, 0.7–2.6, 3.4, 3.5, 3.7, 4.2, 6.8–7.3, 7.5, 7.8–8.2, 8.8–9.0 ppm.

MS: 606.2434

R<sub>f</sub> 0.40 (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 347**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethyl-propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl,  $R_4$  is 5-cyanopyridine-2-yl) [Isomer 3] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the third stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.6, 0.7–2.6, 3.4, 3.5, 3.7, 4.2, 6.8–7.3, 7.5, 7.8–8.2, 8.8–9.0 ppm.

MS: 606.2423

R<sub>f</sub> 0.40 (15% ethyl acetate in dichloromethane)

## **EXAMPLE 348**

N-{3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl} phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl,  $R_4$  is 5-cyanopyridine- 2-yl) [Isomer 4] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the fourth stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–2.7, 3.2, 3.5, 3.6, 3.7, 4.1, 6.8–7.4, 7.5, 7.8–8.2, 8.8 ppm.

HRMS: 606.2429

R<sub>f</sub> 0.40 (15% ethyl acetate in dichloromethane)

## EXAMPLE 349

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl} phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is tert-butyl,  $R_4$  is 1-methylimidazol-4-yl [Isomer 1] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and  $R_3$  is ethyl. The amine is derived from the first stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title

compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.4, 1.7, 2.3–2.7, 3.6, 3.9, 4.1, 6.8–7.5

HRMS: 584.2585

R<sub>c</sub> 0.34 (5% methanol in dichloromethane)

#### **EXAMPLE 350**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or R)- 10 (2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4sulfonamide (Formula WWW-4: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 1-methylimidazol-4-yl [Isomer 2] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the second stereoisomer of Formula WWW-2 to elute from a 20 Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichlo-

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–1.1, 1.3, 1.7, 2.3–2.7, 3.6, 3.9, 4.1, 6.8–7.5

HRMS: 584.2585

 $R_f$  0.34 (5% methanol in dichloromethane)

#### **EXAMPLE 351**

 $N-[3-{1(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } S)-}$ (2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4sulfonamide (Formula WWW-4: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 1-methylimidazol-4-yl [Isomer 3] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the third stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–1.1, 1.3, 1.7, 2.3–2.7, 3.6, 3.9, 4.1, 6.8–7.5

HRMS: 584.2591

R<sub>f</sub> 0.34 (5% methanol in dichloromethane)

### EXAMPLE 352

 $N-[3-\{1(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S \text{ or } R)$ (2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4- 55 sulfonamide (Formula WWW-4: R, is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 1-methylimidazol-4-yl [Isomer 4] Refer to Chart WWW

The title compound is prepared using the general sulfo-Formula WWW-3, where R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl. The amine is derived from the fourth stereoisomer of Formula WWW-2 to elute from a Chiralcel OD chiral HPLC column of Preparation 150. The title compound is obtained as an amorphous solid after flash 65 chromatography on silica using 3% methanol in dichloromethane.

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Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.4, 1.7, 2.3–2.7, 3.6, 3.9, 4.1, 6.8–7.5 ppm.

HRMS: 584.2580

R<sub>f</sub> 0.34 (5% methanol in dichloromethane)

#### **EXAMPLE 353**

 $N-[3-\{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-dihyd$ phenylethyl)-2H-pyran-3-yl)methyl}phenyl]-5cyanopyridine-2-sulfonamide (Formula XXX-5, R<sub>1</sub> is 5-cyanopyridine-2-yl) Refer to Chart XXX

Using the general sulfonylation procedure of Example 252, 64 mg of the amine of formula XXX-4 is reacted with 5-cyanopyridine-2-sulfonyl chloride. Flash chromatography on silica using 2-3% methanol in dichloromethane provides 73.2 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ1.8–2.1, 2.6, 6.9–7.3, 7.9, 8.8 ppm.

HRMS: 594.2068

R<sub>f</sub> 0.40 (3% methanol in dichloromethane)

## **EXAMPLE 354**

N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2phenylethyl)-2H-pyran-3-yl)methyl}phenyl]-5aminopyridine-2-sulfonamide (Formula UUU-6, R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is H). Refer to Chart UUU

Using the general sulfonylation procedure of Example 252, 69 mg of the amine of formula XXX-4 is reacted with 5-nitropyridine-2-sulfonyl chloride. Flash chromatography on silica using 2-3% methanol in dichloromethane provides 107 mg of the intermediate nitro compound of formula UUU-5 (R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is H). Reduction to the amine is accomplished using hydrogen gas and palladium on carbon catalyst. Flash chromatography on silica gel using 4-6% methanol in dichloromethane provides 65.0 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ1.9–2.1, 2.6, 3.5–4.0, 6.7, 6.9–7.3, 7.5, 7.9

HRMS: 584.2215

R<sub>c</sub> 0.24 (5% methanol in dichloromethane)

#### PREPARATION 151

Resolution of N-[3-[1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl] phenyl]carbamic acid, phenylmethyl ester to give 2 enantiomers (Formula WWW-2: R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is t-butyl) Refer to Chart WWW

Inject 40 mg batches of the starting compound onto a 2.1×25 cm (R,R)Whelk-O 1 column (Regis Technologies, Inc.) that is maintained at 30°. The 2 enantiomers elute at about 37 min (Enantiomer 1) and 43 min (Enantiomer 2) using 25% isopropanol and 0.05% acetic acid at 12 mL/min. Fractions are pooled on the basis of results from analysis on a 0.46×25 cm (R,R)Whelk-O 1 column eluted with 30% isopropanol and 0.1% acetic acid (V/V) at 1.0 mL/min. The isomers elute at (Isomer 1) 19.1 and (Isomer 2) 23.0 min respectively.

## **EXAMPLE 355**

nylation procedure of Example 252, using the amine of 60 N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-aminopyridine-2-sulfonamide (Formula WWW-4, R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 5-aminopyridine-2yl) [Enantiomer 1] Refer to Chart WWW

Using the general sulfonylation procedure of Example 252, 73 mg of the amine of Formula WWW-3 ( $R_1$  and  $R_2$  are phenethyl, R<sub>3</sub> is tert-butyl) is reacted with 5-nitropyridine-

2-sulfonyl chloride. The amine used is derived from the first enantiomer of Formula WWW-2 to elute from an (R,R) Whelk-O chiral HPLC column of Preparation 151. Flash chromatography on silica using 5–10% ethyl acetate in dichloromethane provides 94.0 mg of the intermediate nitro compound of formula UUU-5 (R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is tert-butyl). Reduction to the amine is accomplished using hydrogen gas and palladium on carbon catalyst. Flash chromatography on silica gel using 4% methanol in dichloromethane provides 74.8 mg of the title compound as an 10 amorphous white solid.

Physical characteristics are as follows: <sup>1</sup>H NMR δ0.95, 2.0, 2.6, 6.8, 6.9–7.4, 7.5, 7.9 ppm. HRMS: 640.2828 R<sub>f</sub> 0.27 (5% methanol in dichloromethane)

#### **EXAMPLE 356**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-aminopyridine-2-sulfonamide (Formula WWW-4,  $R_1$  and  $R_2$  are phenethyl,  $R_3$  is tert-butyl,  $R_4$  is 5-aminopyridine-2-yl) [Enantiomer 2] Refer to Chart WWW

Using the general sulfonylation procedure of Example 252, 73 mg of the amine of Formula WWW-3 (R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is tert-butyl) is reacted with 5-nitropyridine-2-sulfonyl chloride. The amine used is derived from the second enantiomer of Formula WWW-2 to elute from an (R,R)Whelk-O chiral HPLC column of Preparation 151. Flash chromatography on silica using 5–10% ethyl acetate in dichloromethane provides 91.3 mg of the intermediate nitro compound of formula UUU-5 (R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is tert-butyl). Reduction to the amine is accomplished using hydrogen gas and palladium on carbon catalyst. Flash chromatography on silica gel using 4% methanol in dichloromethane provides 54.3 mg of the title compound as an amorphous white solid.

Physical characteristics are as follows: <sup>1</sup>H NMR 80.95, 2.0, 2.6, 6.8, 6.9-7.4, 7.5, 7.9 ppm. HRMS: 640.2828 R<sub>f</sub> 0.27 (5% methanol in dichloromethane)

## **EXAMPLE 357**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4,  $R_1$  and  $R_2$  are phenethyl,  $R_3$  is tert-butyl,  $R_4$  is 1-methylimidazol-4-yl) [Enantiomer 1] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  and  $R_2$  are phenethyl and  $R_3$  is tert-butyl. The amine is derived from the first enantiomer of Formula WWW-2 to elute from an (R,R)Whelk-O chiral HPLC column of Preparation 151. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows: <sup>1</sup>H NMR 80.98, 2.0, 2.6, 3.6, 3.8, 6.9-7.5 ppm. HRMS: 628.2832 R<sub>f</sub> 0.38 (5% methanol in dichloromethane)

## **EXAMPLE 358**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4,  $R_1$  and  $R_2$  are phenethyl,  $R_3$  is tert-butyl,  $R_4$  is 65 1-methylimidazol-4-yl) [Enantiomer 2] Refer to Chart WWW

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The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where  $R_1$  and  $R_2$  are phenethyl and  $R_3$  is tert-butyl. The amine is derived from the second enantiomer of Formula WWW-2 to elute from an (R,R)Whelk-O chiral HPLC column of Preparation 151. The title compound is obtained as an amorphous solid after flash chromatography on silica using 3% methanol in dichloromethane.

Physical characteristics are as follows: <sup>1</sup>H NMR 80.98, 2.0, 2.6, 3.6, 3.8, 6.9-7.5 ppm. HRMS: 628.2838 R<sub>\*</sub> 0.38 (5% methanol in dichloromethane)

#### **EXAMPLE 359**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4, R<sub>1</sub> and R<sub>2</sub> are phenethyl, R<sub>3</sub> is tert-butyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) [Enantiomer 1] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> and R<sub>2</sub> are phenethyl and R<sub>3</sub> is tert-butyl. The amine is derived from the first enantiomer of Formula WWW-2 to elute from an (R,R)Whelk-O chiral HPLC column of Preparation 151. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

Physical characteristics are as follows:  $^{1}$ H NMR  $\delta 0.87$ , 1.9, 2.6, 6.8–7.4, 7.9, 8.8 ppm. HRMS: 650.2681 R<sub>f</sub> 0.46 (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 360**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4,  $R_1$  and  $R_2$  are phenethyl,  $R_3$  is tert-butyl,  $R_4$  is 5-cyanopyridine-2-yl) [Enantiomer 2] Refer to Chart WWW

The title compound is prepared using the general sulfonylation procedure of Example 252, using the amine of Formula WWW-3, where R<sub>1</sub> and R<sub>2</sub> are phenethyl and R<sub>3</sub> is tert-butyl. The amine is derived from the second enantiomer of Formula WWW-2 to elute from an (R,R)Whelk-O chiral HPLC column of Preparation 151. The title compound is obtained as an amorphous solid after flash chromatography on silica using 10% ethyl acetate in dichloromethane.

<sup>1</sup>H NMR δ0.87, 1.9, 2.6, 6.8–7.4, 7.9, 8.8 ppm. HRMS: 650.2681 R<sub>f</sub> 0.46 (15% ethyl acetate in dichloromethane)

Physical characteristics are as follows:

#### PREPARATION 152

Resolution of N-[3-[1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl]phenyl]carbamic acid, phenylmethyl ester to give 2 isomers (Formula WWW-2:  $R_1$  and  $R_2$  are propyl,  $R_3$  is ethyl) Refer to Chart WWW

Samples of the starting compound are injected onto a 2.1×25 cm Chiralcel OD column and eluted with 20% isopropanol (V/V) in hexane at 10 mL/min. The material eluting near 19.1 minutes is one isomer (Enantiomer 1) and that eluting near 37.7 minutes is another isomer (Enantiomer 2). The pools are concentrated separately on a rotary evaporator (ca. 30 mm, bath at 50° maximum) to give white solids.

### **EXAMPLE 361**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4,  $R_1$  and  $R_2$  are propyl,  $R_3$  is ethyl,  $R_4$  is 5-cyanopyridine-2-yl) [Enantiomer 1] Refer to Chart WWW

Following procedures analogous to those described above, but using Enantiomer 1 of Preparation 152, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.2–2.2, 3.90, 6.9–7.2, 8.0, 8.15, 8.9 5

HRMS: 497.1984

R<sub>f</sub> 0.38 (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 362**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4, R<sub>1</sub> and R<sub>2</sub> are propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) [Enantiomer 2] Refer to Chart WWW

Following procedures analogous to those described above, but using Enantiomer 2 of Preparation 152, the title compound is obtained.

Physical characteristics are as follows:

ppm.

HRMS: 497.1980

R<sub>c</sub> 0.38 (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 363**

 $N-[3-\{1(R \ or \ S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dihydro-2-o$ dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-aminopyridine-2-sulfonamide (Formula WWW-4, R<sub>1</sub> and R<sub>2</sub> are propyl, R<sub>3</sub> is ethyl, R4 is 5-aminopyridine-2-yl) [Enantiomer 1] Refer to Chart WWW

Following procedures analogous to those described 30 above, but using Enantiomer 1 of Preparation 152, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–0.9, 1.2–2.2, 3.8, 6.8–7.2, 7.5, 7.9 ppm. HRMS: 487.2122

R<sub>f</sub> 0.28 (5% methanol in dichloromethane)

## **EXAMPLE 364**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-aminopyridine-  $^{40}$ 2-sulfonamide (Formula WWW-4,  $R_1$  and  $R_2$  are propyl,  $R_3$ is ethyl, R4 is 5-aminopyridine-2-yl) [Enantiomer 2] Refer to Chart WWW

Following procedures analogous to those described above, but using Enantiomer 2 of Preparation 152, the title 45 compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.7–0.9, 1.2–2.2, 3.8, 6.8–7.2, 7.5, 7.9 ppm. HRMS: 487.2140

R<sub>c</sub> 0.28 (5% methanol in dichloromethane)

### PREPARATION 153

Resolution of N-[3-[1-(4-Hydroxy-5,6-dihydro-2-oxo-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)propyl]phenyl]carbamic acid, phenylmethyl ester to give 4 isomers 55 (Formula WWW-2: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, and R<sub>3</sub> is ethyl) Refer to Chart WWW

The enantiomers are defined by elution order from System D. HPLC System D consists of a 0.46×25 cm Chiralcel OD-H column (Chiral Technologies, Inc.) with 20% isopro- 60 panol and 0.05% trifluoroacetic acid in hexane (V/V) pumped at 0.5 mL/min. The retention times in this system are (Isomer 1) 21.6, (Isomer 2) 34.5, (Isomer 3) 55.2 and (Isomer 4) 66.6 min.

Separate the enantiomers on a 2.1×25 cm Chiralcel OD 65 column (Chiral Technologies, Inc.). Aliquots are injected and the enantiomers eluted with 17.5% isopropanol in

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hexane (V/V) at 10 mL/min. Fractions eluting near 24.6, 42.9, 66.3 and 77.4 min are pooled appropriately after assay with System D. In order of elution, the four isomers are designated Isomers 1-4, respectively.

In all cases, whenever solvent is stripped from a pool the following protocol is used: Solvent is removed from pools of fractions on a rotary evaporator with house vacuum (ca. 30 mm Hg) and a water bath set at 45°±5°. If acetic acid is present in the solvent, add ca. 10 mL of toluene/L of pool before the flask goes dry. Residues are then washed into tared flasks using methylene chloride and the solvent is stripped as above. Final solvent removal is accomplished at ambient temperature, 1 mmHg pressure for 2-24 hours before weighing.

### **EXAMPLE 365**

 $N-[3-{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R \text{ or } S)-}$ (2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl,  $^{1}$ H NMR  $\delta 0.8-1.0$ , 1.2-2.2, 3.90, 6.9-7.2, 8.0, 8.15, 8.9  $_{20}$   $_{1}$   $_{20}$ www

> Following procedures analogous to those described above, but using Isomer 1 of Preparation 153, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR  $\delta 0.8$ –1.0, 1.3, 1.6–2.2, 2.5, 3.9, 6.8–7.3, 7.9–8.1, 8.9 ppm.

HRMS: 578.2120

R<sub>f</sub> 0.35 (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 366**

 $N-[3-\{1(R \text{ or } S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S \text{ or } R)-(4-Hydroxy-5,6-dihydro-2-oxo-6)-(4-Hydr$ (2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is ethyl, 35 R<sub>4</sub> is 5-cyanopyridine-2-yl) [Isomer 2] Refer to Chart www

Following procedures analogous to those described above, but using Isomer 2 of Preparation 153, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.3, 1.6–2.2, 2.5, 3.9, 6.8–7.3, 7.9–8.1,

HRMS: 578.2120

 $R_f 0.35$  (15% ethyl acetate in dichloromethane)

#### **EXAMPLE 367**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide (Formula WWW-4: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) [Isomer 3] Refer to Chart

Following procedures analogous to those described above, but using Isomer 3 of Preparation 153, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.3, 1.6–2.2, 2.5, 3.9, 6.8–7.3, 7.9–8.1, 8.9 ppm.

HRMS: 578.2126

R<sub>f</sub> 0.35 (15% ethyl acetate in dichloromethane)

## **EXAMPLE 367A**

N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S or R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine- 2-sulfonamide (Formula WWW-4: R<sub>1</sub> is 4-fluorophenethyl, R<sub>2</sub> is propyl, R<sub>3</sub> is ethyl, R<sub>4</sub> is 5-cyanopyridine-2-yl) [Isomer 4] Refer to Chart

Following procedures analogous to those described above, but using Isomer 4 of Preparation 153, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.3, 1.6–2.2, 2.5, 3.9, 6.8–7.3, 7.9–8.1, 5 8.9 ppm.

HRMS: 578.2126

R<sub>f</sub> 0.35 (15% ethyl acetate in dichloromethane)

## **EXAMPLE 368**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 1-methylimidazol-4-yl) [Isomer 1] Refer to Chart WWW

Following procedures analogous to those described above, but using Isomer 1 of Preparation 153, the title compound are as follows:

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.3, 1.6–2.2, 2.6, 3.63, 4.0, 6.9–7.5

HRMS: 556.2265

R<sub>f</sub> 0.29 (5% methanol in dichloromethane)

#### **EXAMPLE 369**

N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-aminopyridine-2-sulfonamide (Formula WWW-4:  $R_1$  is 4-fluorophenethyl,  $R_2$  is propyl,  $R_3$  is ethyl,  $R_4$  is 5-aminopyridine-2-yl) [Isomer 1] Refer to Chart 30 WWW

Following procedures analogous to those described above, but using Isomer 1 of Preparation 153, the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR δ0.8–1.0, 1.3, 1.6–2.2, 2.5, 3.9, 6.8–7.2, 7.5, 7.9 ppm.

HRMS: 568.2271

R<sub>f</sub> 0.27 (5% methanol in dichloromethane)

### PREPARATION 154

Hexahydro-2H-1-benzopyran-2,4(3H)-dione (Formula DDDD-2, wherein n is 1) Refer to Chart DDDD

A solution of 0.42 g of platinum oxide and 1.66 g of the compound of formula DDDD-1 wherein n is 1 in 100 mL of acetic acid is placed on a Parr hydrogenation apparatus under an initial pressure of 50 psi of hydrogen for 1.5 h. The reaction mixture is then filtered through Celite and concentrated in vacuo to give a beige solid. The crude material is purified by flash column chromatography on silica gel 60 (230–400 mesh) eluting with 0-5% methanol in chloroform to give 0.94 g of the title product as a white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 84.84–4.80, 3.54, 3.40, 2.60–2.53, 2.08–2.02, 1.79–1.65, 1.62–1.54, 1.44–1.40 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ203.0, 167.4, 74.3, 47.7, 45.6, 29.1, <sup>55</sup> 23.5, 23.2, 19.7 ppm.

IR (mineral oil) 3092, 2768, 2714, 2695, 2662, 1657, 1614, 1577, 1444, 1352, 1345, 1340, 1323, 1308, 1295, 1287, 1260, 1244, 1211, 1188, 1057, 1004, 938, 909, 890, 843, 832, 600 cm<sup>-1</sup>.

EI-MS: [M+]=168.

Anal. found: C, 64.16; H, 7.16.

## PREPARATION 155

4a,5,6,7,8,8a-Hexahydro-4-hydroxy-3-[1-(3-nitrophenyl) 65 propyl]-2H-1-benzopyran-2-one (Formula DDDD-4, wherein n is 1 and  $R_1$  is ethyl) Refer to Chart DDDD

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A solution of 3.17 g of aluminum trichloride in 30 mL of tetrahydrofuran is added to a solution of 2.00 g of the title compound of Preparation 154 and 1.82 g of 3-nitrobenzaldehyde in 20 mL of tetrahydrofuran. The resulting mixture is then stirred at room temperature for 2.5 h, at which time, 7.28 g of sodium carbonate decahydrate is added, and the reaction mixture is stirred an additional 20 min. The mixture is then dried over magnesium sulfate, filtered through Celite, and concentrated in vacuo to yield 10 6.05 g of a yellow gum. This crude material is immediately dissolved in 50 mL of tetrahydrofuran containing 0.73 g of cuprous bromide-dimethyl sulfide complex, and 13.1 mL of a 1.0M solution of triethyl aluminum in hexanes are added to the reaction mixture. After stirring at room temperature for 1 h, the reaction is quenched by the addition of water, and the resulting mixture is partitioned between ether and water. The organic layer is separated, washed with brine, and concentrated in vacuo to produce 4.0 g of a yellow oil. The crude material is purified by flash column chromatography eluting with 10-50% ethyl acetate in hexanes to yield 0.63 g of the title product as a yellow foam.

Physical characteristics are as follows:

MP 86°-91° C.

IR (mineral oil) 3085, 1635, 1569, 1528, 1448, 1394, 25 1365, 1349, 1325, 1307, 1288, 1270, 1251, 1244 cm $^{-1}$ .

#### **EXAMPLE 370**

5-Cyano-N-[3-1-(4a,5,6,7,8,8a-hexahydro-4-hydroxy-2-oxo-2H-1-benzopyran-3-yl)propyl]phenyl]-2-pyridinesulfonamide (Formula DDDD-7, wherein n is 1,  $R_1$  is ethyl, and  $R_2$  is 5-cyano-2-pyridyl) Refer to Chart DDDD

A solution of 0.63 g of the title compound of Preparation 155 in 50 mL of ethanol with 0.3 g of 10% palladium on carbon is placed on a Parr hydrogenation apparatus at an initial pressure of 50 psi of hydrogen for 3 h. The reaction mixture is then filtered through Celite and concentrated in vacuo to give 0.519 g of crude intermediate. 0.25 g of this intermediate is immediately dissolved in 5 mL of methylene chloride, and 0.168 g of 5-cyano-2-pyridylsulfonyl chloride and 0.134 mL of pyridine are added to the solution. The resulting mixture is stirred at room temperature for 18 h. The reaction mixture is then purified by flash column chromatography on silica gel 60 (230-400 mesh) eluting with 0-2.5% methanol in chloroform to give 0.164 g of the title product as a white foam.

Physical characteristics are as follows:

MP 122°-125° C. HRMS found: 468.1611

# **EXAMPLE 371**

 $^{50}$  4-Cyano-N-[3-1-(4a,5,6,7,8,8a-hexahydro-4-hydroxy-2-oxo-2H-1-benzopyran-3-yl)propyl]phenyl]-benzenesulfonamide (Formula DDDD-7, wherein n is 1,  $R_{\rm 1}$  is ethyl, and  $R_{\rm 2}$  is 4-cyanophenyl) Refer to Chart DDDD

Following the general procedure of Example 370, and making non-critical variations, but substituting 4-cyanophenylsulfonyl chloride for 5-cyano-2-pyridylsulfonyl chloride, 0.236 g of the title compound is obtained as white foam.

Physical characteristics are as follows:

MP 127°-130° C.

HRMS found: 466.1583.

#### PREPARATION 156

4-Hexahydro-cyclohepta[b]pyran-2,4(3H,4aH)-dione (Formula DDDD-2, wherein n is 2) Refer to Chart DDDD Following the general procedure of Preparation 154, and making non-critical variations, but substituting the cyclo-

heptylpyranone of Formula DDDD-1 wherein n is 2 for the cyclohexylpyranone of Formula DDDD-1 wherein n is 1, 0.337 g of the title compound is obtained as white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR CDCl<sub>3</sub>) 84.97–4.91, 3.52, 3.42, 2.64–2.58, 5 2.22–2.11, 2.01–1.72, 1.59–1.36 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ203.0, 167.2, 78.0, 52.1, 46.5, 32.1,

28.6, 27.1, 25.7, 21.3 ppm.

IR (mineral oil) 3074, 2791, 2755, 2736, 2687, 2637, 2608, 2585, 1655, 1625, 1586, 1500, 1480, 1443, 1333, 10 1324, 1293 (s), 1265, 1254, 1240, 1222, 1196, 1173, 1082, 1053, 1016, 909, 889, 832, 611 cm<sup>-1</sup>.

EI-MS: [M+]=182.

Anal. found: C, 66.16; H, 7.90.

#### PREPARATION 157

5,6,7,8,9,9a-Hexahydro-4-hydroxy-3-[1-(3-nitrophenyl)-propyl]-cyclohepta[b]pyran-2(4aH)-one (Formula DDDD-4, wherein n is 2 and R<sub>1</sub> is ethyl) Refer to Chart DDDD

Following the general procedure of Preparation 155, and making non-critical variations, but substituting the title compound of Preparation 156 for the title compound of Preparation 154, 2.5 g of the title compound is obtained as a yellow foam.

Physical characteristics are as follows:

MP 75°-78° C.

IR (mineral oil) 3071, 2667, 1638, 1528, 1395, 1350, 1305, 1276, 1250, 1143, 1130, 1120, 1100, 1066, 782, 764, 741, 697,  $685 \text{ cm}^{-1}$ .

HRMS found: 345.1590.

Anal. found: C, 58.74; H, 5.63; N, 3.48.

#### **EXAMPLE 372**

5-Cyano-N- $\{3-[1-(2,4a,5,6,7,8,9,9a-\text{octahydro-}4-\text{hydroxy-}2-\text{oxocyclohepta}[b]pyran-3-yl)propyl]phenyl]-2-pyridinesulfonamide (Formula DDDD-7, wherein n is 2, <math>R_1$  is ethyl, and  $R_2$  is 5-cyano-2-pyridyl) Refer to Chart DDDD

Following the general procedure of Example 370, and making non-critical variations, but substituting the title compound of Preparation 157 for the title compound of Preparation 155, 0.206 g of the title compound is obtained as a white foam.

Physical characteristics are as follows:

MP 163°-166° C.

IR (mineral oil) 3352, 3128, 3100, 3073, 3029, 1760, 1726, 1641, 1608, 1593, 1584, 1411, 1397, 1355, 1295, 1282, 1242, 1207, 1173, 1125, 1106, 1086, 1074, 1028, 974, 967, 721, 701, 645, 638 cm<sup>-1</sup>.

HRMS found: 481.1693.

## PREPARATION 158

Octahydro-2H-cycloocta[b]pyran-2,4(3H)-dione (Formula DDDD-2, wherein n is 3) Refer to Chart DDDD

Following the general procedure of Preparation 154, and making non-critical variations, but substituting the cyclooctylpyranone of Formula DDDD-1 wherein n=3 for the 55 cycloheptylpyranone of Formula DDDD-1 wherein n=2, 1.72 g of the title compound is obtained as a white solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 84.84–4.78, 3.61, 3.40, 2.75–2.70, 2.14–1.97, 1.90–1.72, 1.68–1.44 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ204.2, 167.2, 78.2, 49.5, 46.1, 28.5, 27.3, 26.2, 24.7, 23.9, 22.1 ppm.

IR (mineral oil) 2659, 2617, 1650, 1612, 1579, 1444, 1356, 1332, 1307, 1287, 1265, 1244, 1227, 1209, 1041, 1035, 1003, 962, 946, 860, 832, 824 cm<sup>-1</sup>.

HRMS found: 196.1100.

Anal. found: C, 67.06; H, 8.23.

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## PREPARATION 159

3-[2,2-Dimethyl-1-(3-nitrophenyl)propyl]-4a,5,6,7,8,9,10, 10a-octahydro-4-hydroxy-2H-cycloocta[b]pyran-2-one (Formula DDDD-4, wherein n is 3 and  $R_1$  is t-butyl) Refer to Chart DDDD

A solution of 1.36 g of aluminum trichloride in 30 mL of tetrahydrofuran is added to a solution of 1.0 g of the title compound of Preparation 158 and 0.77 g of 3-nitrobenzaldehyde in 20 mL of tetrahydrofuran. The resulting mixture is then stirred at room temperature for 2.3 h, at which time, 3.06 g of sodium carbonate decahydrate is added, and the reaction mixture is stirred an additional 15 min. The mixture is then dried over magnesium sulfate, filtered through Celite, and concentrated in vacuo to yield a yellow foam. This crude intermediate is immediately dissolved in 5 mL of tetrahydrofuran for use in the second step.

A dry flask is charged with 0.82 g of activated zinc, 3 mL of tetrahydrofuran, 0.035 mL of dibromoethane, and 0.21 mL of a 1M solution of trimethylsilyl chloride in tetrahydrofuran. After the addition of each reagent the mixture is sonicated for 15 m at 45° C. The mixture is diluted further by the addition of 2 mL tetrahydrofuran and 1.32 mL of t-butyl iodide is added dropwise. The resulting mixture is sonicated for 3 h at 45° C. A separate mixture of 0.85 g of copper(I) cyanide and 0.80 g of lithium chloride in 4 mL of tetrahydrofuran is stirred at room temperature for 1 h until almost homogeneous and cooled to -30° C. The organozinc solution is then added via cannula to the copper cyanide solution and the resulting mixture is allowed to warm to 0° C. and to stir for 15 min. The reaction mixture is then cooled to -78° C., and the solution of crude intermediate prepared above is added. After stirring for 20 min at -78° C. and 30 min at 0° C., the reaction is quenched with a saturated solution of aqueous ammonium chloride and diluted with an additional 60 mL of tetrahydrofuran. The organic layer is separated, washed with water, and concentrated in vacuo to give 2.17 g of an orange foam. The crude material is then purified by flash column chromatography eluting with 10-30% ethyl acetate in hexanes followed by recrystallization in methylene chloride/hexanes to yield 0.60 g of the title product as a yellow solid.

Physical characteristics are as follows:

MP 158°-161° C.

IR (mineral oil) 3077, 2646, 1632, 1599, 1529, 1477, 1450, 1396, 1357, 1349, 1334, 1317, 1283, 1273, 1252, 1232, 1217, 1205, 1181 cm<sup>-1</sup>.

## **EXAMPLE 373**

5-Cyano-N-[3-[2,2-dimethyl-1-(4a,5,6,7,8,9,10,10a-octahydro-4-hydroxy-2-oxo-2H-cycloocta[b]pyran-3-yl) propyl]phenyl]-2-pyridinesulfonamide (Formula DDDD-7, wherein n is 2,  $R_1$  is t-butyl, and  $R_2$  is 5-cyano-2-pyridyl) Refer to Chart DDDD

Following the general procedure of Example 370, and making non-critical variations, but substituting the title compound of Preparation 159 for the title compound of Preparation 157, 0.034 g of the title compound is obtained as white crystals.

Physical characteristics are as follows:

MP 182°-185° C.

IR (mineral oil) 3246, 3121, 3098, 2615, 1655, 1633, 65 1607, 1585, 1575, 1491, 1411, 1395, 1354, 1335, 1322, 1311, 1298, 1281, 1275, 1262, 1255, 1233, 1206, 1178, 1121, 1109, 1028, 977, 702, 657, 646, 635, 605 cm<sup>-1</sup>.

Anal. found: C, 63.86; H, 6.41; N, 7.82.

## PREPARATION 160

 $(3(3R),4S)-3-[2-[1-[3-[bis(phenylmethyl)amino]phenyl]-propyl]-5-hydroxy-1,3-dioxo-5-(2-phenylethyl)octyl]-4-phenyl-2-oxazolidinone (Formula W-10 wherein <math>R_1$  is 2-phenylethyl) Refer to Chart W

To 100 mL of methylene chloride is added 5.0 g of the title compound of Preparation 95 (W-8) and the resulting solution cooled to -78° C. under an atmosphere of nitrogen. To that solution is added 1.0 mL of TiCl<sub>4</sub> and 1.63 mL of disopropylethylamine, and the resulting solution is stirred for 1 hour. Then, 3.30 g of 1-phenyl-3-hexanone is added, and the reaction temperature raised to 0° C. for 2.5 hours. The reaction is then quenched by the addition of a saturated ammonium chloride solution, and the mixture is extracted with methylene chloride. The organic extract is washed with saturated sodium bicarbonate solution and evaporated in vacuo to yield 9.7 g of a yellow oil. Column chromatography on 900 g silica (elution with 10% hexane-methylene chloride, 100% methylene chloride) affords 3.30 g of the title compound as a yellow foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.33–7.23, 7.14, 7.04, 6.61–6.50, 5.45, 5.22, 4.71, 4.60, 4.48, 4.26, 3.33, 3.15–3.03, 2.58, 2.47–2.31, 1.93, 1.40–1.28, 1.24–1.13, 1.11–0.96, 0.88–0.77, 0.62–0.57 ppm.

MP 121°-126° C.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) 8167.2, 167.1, 153.7, 142.6, 141.0, 138.2, 138.1, 129.6, 129.5, 129.2, 128.9, 128.8, 128.6, 128.4, 128.3, 127.0, 125.8, 125.6, 125.6, 73.1, 70.0, 69.9, 63.9, 57.9, 54.8, 54.7, 51.5, 51.4, 48.3, 41.3, 41.0, 40.8, 40.5, 29.8, 29.6, 27.1, 26.9, 16.8, 16.6, 14.6, 11.7, 11.6 ppm.

IR (mineral oil) 3525, 3061, 3026, 1777, 1720, 1690, 1601, 1495, 1361, 1335, 1238, 1199, 1104, 735, 698 cm<sup>-1</sup>. EI-MS: [M+]=736.

Anal. found: C, 78.03; H, 7.11; N, 3.79.

## PREPARATION 161

(3S)-3-[1-[3-Bis(phenylmethyl)amino]phenyl]propyl]-5,6-40 dihydro-4-hydroxy-6-(2-phenylethyl-6-propyl-pyran-2-one (Formula W-11 wherein  $R_1$  is 2-phenylethyl) Refer to Chart

To 5 mL of dry tetrahydrofuran is added 2.7 g of the title compound of Preparation 160 and the resulting solution is 45 cooled to 0° C. under an atmosphere of nitrogen. To that solution is added 0.45 mL of a 1M solution of potassium t-butoxide in tetrahydrofuran. The reaction mixture is then warmed to 20° C. and stirred for 2 hours. The reaction is quenched with saturated ammonium chloride and extracted 50 with ethyl acetate. The organic layer is washed with water, dried and evaporated in vacuo to yield 0.28 g of a yellow oil. Column chromatography on 80 g of silica gel (elution with 10–30% acetone/hexane) affords 0.195 g of a yellow foam. Crystallization from ethyl acetate/hexane yields 0.146 g of 55 the title compound.

Physical characteristics are as follows:

MP 128°-131° C

<sup>1</sup>H NMR (CDCl<sub>3</sub>) δ7.35–7.12, 6.73–6.64, 5.84, 4.73–4.57, 4.12, 2.69–2.61, 2.38–2.20, 1.95–1.65, 60 1.41–1.32, 0.98–0.87 ppm.

<sup>13</sup>C NMR (CDCl<sub>3</sub>) 8204.1, 204.0, 171.7, 171.4, 169.6, 140.9, 140.8, 140.6, 140.5, 140.4, 139.9, 139.8, 138.3, 129.7, 129.6, 129.5, 128.9, 128.6, 128.5, 128.4, 128.2, 128.1, 127.1, 126.9, 126.8, 126.7, 126.5, 126.4, 126.3, 126.2, 126.0, 125.9, 116.8, 112.6, 112.5, 112.4, 112.3, 112.2, 112.1, 112.0, 82.0, 81.9, 81.8, 80.4, 80.3, 58.6, 58.5, 54.5,

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51.4, 50.4, 50.1, 49.9, 47.8, 47.4, 47.0, 46.6, 43.0, 42.9, 42.2, 41.9, 40.2, 40.1, 40.0, 39.2, 29.8, 29.7, 29.6, 29.1, 29.0, 26.8, 26.7, 24.7, 24.6, 24.3, 16.9, 16.5, 14.0, 12.3 ppm.

IR (mineral oil) 3023, 1637, 1599, 1584, 1575, 1494, 1347, 1300, 1257, 1243, 1234, 920, 731, 704, 695 cm<sup>-1</sup>.

EI-MS: [M+]=573.

Anal. found: C, 81.53; H, 7.82; N, 2.34. [a]<sub>D</sub> (CHCl<sub>3</sub>)=-83°

### PREPARATION 162

(3S)-3-[1-(3-Aminophenyl)propyl]-5,6-dihydro-4-hydroxy-6-(2-phenylethyl)-6-propyl-pyran-2-one (Formula W-12 wherein  $R_1$  is 2-phenylethyl) Refer to Chart W

0.63 g of the title compound of Preparation 161 is dissolved in 45 mL of ethyl acetate and 15 mL of methanol. To that solution is added 0.47 g of 10% Pd/C, and the resulting mixture is hydrogenated at 50 psi for 2.5 hours. The reaction is then filtered through celite and concentrated in vacuo to yield 0.466 g of an off-white foam. Column chromatography on 80 g of silica gel (elution with 20-50% ethyl acetate-hexane) affords 0.389 g of the title compound as an off-white solid.

Physical characteristics are as follows:

MÝ 155°-159° C.

<sup>1</sup>H NMR (CD<sub>3</sub>OD) δ7.26–7.20, 7.15–7.04, 6.95, 6.81, 6.74, 6.54–6.51, 3.98–3.91, 2.68–2.54, 2.25–2.17, 2.02–1.67, 1.43–1.28, 0.99–0.87 ppm.

<sup>13</sup>C NMR (CD<sub>3</sub>OD) δ171.2, 171.0, 148.5, 148.2, 143.9, 130.4, 130.2, 127.8, 120.8, 117.8, 115.3, 107.4, 82.7, 44.6, 44.4, 41.8, 41.7, 41.5, 38.4, 31.8, 26.8, 26.7, 18.8, 15.6, 14.3 ppm.

IR (mineral oil) 3085, 3061, 3026, 1617, 1605, 1495, 1314, 1258, 1168, 1119, 1065, 1030, 923, 776, 699 cm<sup>-1</sup>. EI-MS: [M+]=393.

Anal. found: C, 76.13; H, 8.16; N, 3.37.  $[a]_D$  (MeOH)=-41°

## **EXAMPLE 374**

N-[3-[1-(S)-[5,6,-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide (Formula W-13 wherein R<sub>1</sub> is 2-phenylethyl) Refer to Chart W

To a solution of 0.200 g of the title compound of Preparation 162 in 5 mL of methylene chloride is added 0.12 mL of pyridine. The resulting mixture is cooled to 0° C. and 0.132 g of 5-trifluoromethylpyridine-2-sulfonyl chloride is added. The reaction mixture is then stirred at room temperature for 1.5 h, concentrated in vacuo, and partitioned between ethyl acetate and water. The organic layer is concentrated in vacuo to 0.39 g of pink oil. Column chromatography on 50 g silica gel (elution with 20-50% ethyl acetate/hexane) affords 0.252 g of title compound as a white foam.

MP 170°-173° C.

<sup>1</sup>H NMR (CD<sub>3</sub>OD) δ8.95–8.92, 8.23–8.16, 8.04–8.00, 7.25–6.90, 4.86, 3.98–3.90, 3.31, 3.30, 2.69–2.46, 2.18–2.09, 1.96–1.65, 1.41–1.28, 0.99–0.81 ppm.

<sup>13</sup>C NMR (CD<sub>3</sub>OD) δ167.3, 147.7, 147.5, 142.9, 142.8, 137.7, 137.0, 129.5, 129.2, 126.9, 126.2, 126.1, 124.1, 122.6, 122.5, 120.3, 120.2, 81.8, 81.7, 43.6, 43.2, 40.9, 40.5, 37.5, 30.9, 25.8, 25.6, 17.9, 14.7, 13.3, 13.2 ppm.

IR (mineral oil) 3087, 3027, 1642, 1606, 1595, 1327, 1260, 1173, 1142, 1110, 1074, 1016, 720, 700, 613 cm<sup>-1</sup>. FAB-MS: [M+H]=603.

Anal. found: C, 61.79; H, 5.86; N, 4.48; S, 5.16. [a]<sub>D</sub> (MeOH)=-310.

## PREPARATION 163

(3S,6R)-3-[1-[3-bis(phenylmethyl)amino]phenyl]propyl]-5, 6-dihydro-4-hydroxy-6-(2-phenylethyl-6-propyl-pyran-2one (Formula FFF-2) Refer to Chart FFF

The title compound of Preparation 161 is chromatographed on a 5.1×30 cm Cyclobond I 2000 column in an ice bath at 90 mg per injection using an automated chromatographic system and a mobile phase of acetonitrile containing 0.1% diethylamine and 0.05% glacial acetic acid (v/v). The eluant is monitored at 260 nm, the flow rate is 45 mL/min, and appropriate fractions from multiple injections are combined and concentrated in vacuo to give 0.300 g of a dark oil. The oil is partitioned between ethyl acetate, saturated aqueous sodium bicarbonate solution, and water. The organic 10 layer is separated and concentrated in vacuo. Column chromatography on 50 g of silica gel (elution with 10–20% acetone/hexane) affords 0.22 g of the title compound of the compound as a colorless oil.

Physical characteristics are as follows: The retention time of the title compound is 57 min.

## PREPARATION 164

(3S,6S)-3-[1-[3-bis(phenylmethyl)amino]phenyl]propyl]-5, 6-dihydro-4-hydroxy-6-(2-phenylethyl-6-propyl-pyran-2one (Formula FFF-3) Refer to Chart FFF

The title compound of Preparation 161 is separated as described in Preparation 163 above. Further purification as described in Preparation 163 affords 0.117 g of the title compound as a colorless oil.

Physical characteristics are as follows:

The retention time of the title compound is 66 min.

## PREPARATION 165

(3S,6R)-3-[1-(3-aminophenyl)propyl]-5,6-dihydro-4-hydroxy-6-(2-phenylethyl)-6-propyl-pyran-2-one (Formula FFF-4) Refer to Chart FFF

Following the general procedure of Preparation 162, and making non-critical variations, but substituting the title product of Preparation 163 for the title product of Preparation 161, 0.022 g of the title compound is obtained.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD)  $\delta$ 7.25–7.18, 7.15–7.12, 7.07–7.05, 6.97, 6.82-, 6.76–6.71, 6.53, 4.00–3.92, 2.67–2.54, 2.29–2.15, 2.06–1.92, 1.90–1.62, 1.46–1.28, 0.97–0.88 ppm.

## **EXAMPLE 375**

(3S,6R)-N-[3-[1-[5,6,-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide (Formula FFF-5) <sup>45</sup> Refer to Chart FFF

Following the general procedure of Example 374, and making non-critical variations, but substituting the title product of Preparation 165 for the title product of Preparation 162, 0.024 g of the title compound is obtained as a white foam.

Physical characteristics are as follows:

MP 156°-159° C.

<sup>1</sup>H NMR (CD<sub>3</sub>OD) δ8.90, 8.20–8.17, 8.02–7.99, 7.28–6.88, 4.00–3.90, 2.71–2.46, 2.20–2.10, 1.98–1.67, <sup>55</sup> 1.41–1.28, 0.98–0.81 ppm.

## **EXAMPLE 376**

(3S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6-dipropyl-2H-pyran- 3-yl)propyl]phenyl]-5-(trifluoromethyl)-2- 60 pyridinesulfonamide (Formula W-13) Refer to Chart W

The title compound of Preparation 99 (formula W-12) 182 mg is dissolved in 5 mL of methylene chloride and 133  $\mu$ L of pyridine added. The reaction is cooled to 0° C. and 142 mg of 5-trifluoromethyl-2-pyridinesulfonylchloride added. 65 The reaction is stirred for 30 minutes and the methylene chloride is evaporated and the resulting material diluted with

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ethyl acetate. The organic solution is washed with water, brine and then dried over sodium sulfate. Evaporation of solvent gives 580 mg of crude product. Silica gel chromatography (50 g) eluting with 50% ethyl acetate/hexane affords 211 mg of the desired product as a white foam.

Physical characteristics are as follows: Anal. found: C, 57.80; H, 5.95; N, 5.01; S, 5.64  $[\alpha]_D$  (18.094 mg/2 mL CHCl<sub>3</sub>)=-30°

#### **EXAMPLE 377**

(3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide (Formula X-13) Refer to Chart X

The title compound of Preparation 107 (formula X-12) 170 mg is dissolved in 5 mL of methylene chloride and 136 µL of pyridine added. The reaction is cooled to 0° C. and 132 mg of 5-trifluoromethyl-2-pyridinesulfonylchloride is added. The reaction is stirred for 30 minutes and the methylene chloride is evaporated and the resulting material diluted with ethyl acetate. The organic solution is washed with water, brine and then dried over sodium sulfate. Evaporation of solvent gives crude product which is chromatographed over 50 g of silica gel eluting with 50% ethyl acetate/hexane affords 225 mg of the desired product as a white foam.

Physical characteristics are as follows:  $[\alpha]_D$  (mg/2 mL CHCl<sub>3</sub>)=+29°

## PREPARATION 166

(3S)(4R) 3-[2-[1-[3-[Bis(phenylmethyl)amino]phenyl]-propyl]-5-hydroxy-1,3-dioxo-5-phenethyloctyl]-4-phenyl-2-oxazolidinone (Formula X-10 where R<sub>1</sub> is phenethyl) Refer to Chart X

To 1.12 g of the title compound of Preparation 104 is added 20 mL of methylene chloride and the resulting solution cooled to  $-78^{\circ}$  C. To that solution is added 237  $\mu$ L of TiCl<sub>4</sub> followed by 400  $\mu$ L of diisopropylethylamine and the resulting solution is stirred at  $-78^{\circ}$  C. for 1 hour. To the aforementioned solution is added 776  $\mu$ L of 1-phenyl-3-hexanone and stirring continued at  $-40^{\circ}$  C. for 40 minutes and then the temperature is raised to  $-10^{\circ}$  C. for 1.5 hours. The reaction is quenched with the addition of a saturated ammonium chloride solution, then extraction with methylene chloride and evaporation of the organic extracts. The crude material is chromatographed over 200 g of silica gel eluting with 10% hexane/methylene chloride to afford 870 mg of the title compound.

Physical characteristics are as follows:

IR (mineral oil) 2956, 2926, 2854, 1777, 1600, 1494, 1452, 698 cm<sup>-</sup>1.

 $[\alpha]_D$  (16.578 mg in CHCl<sub>3</sub>)=+4° Mass Spectrum: molecular ion at 736. Anal. found. C, 78.00; H, 7.14; N, 3.61.

## PREPARATION 167

(3R) 3-[1-[3-{Bis(phenylmethyl)amino]phenyl]propyl]-5,6-dihydro-4-hydroxy-6-phenethyl-6-propyl-2H-pyran-2-one (Formula X-11 where  $R_1$  is phenethyl) Refer to Chart X

The compound of Preparation 166 (750 mg) is added to 5 mL of dry THF and potassium tert. butoxide (1.0M in THF; 1.2 mL) is added. The reaction is stirred at 20° C. for 30 minutes and then quenched by the addition of a saturated ammonium chloride solution. The reaction is extracted with ethyl acetate, the organic extracts washed with water and brine and finally evaporated to afford the crude product. Silica gel chromatography over 100 g of silica gel eluting with 15% ethyl acetate/hexane affords 511 mg of the title product.

Physical characteristics are as follows:

IR (mineral oil) 2956, 2855, 1628, 1599, 1577, 1494, 1385, 1364, 697 cm<sup>-</sup>1.

Anal. found: C, 81.30; H, 7.68; N, 2.30 Mass spectrum: molecular ion at 573.  $[\alpha]_D$  (18.116 mg/2 mL CH<sub>3</sub>OH)=+38°

## PREPARATION 168

(3R) 3-[1-[3-aminophenyl]propyl]-5,6-dihydro-4-hydroxy-6-phenethyl-6-propyl-2H-pyran-2-one (Formula X-12 where R<sub>1</sub> is phenethyl) Refer to Chart X

The compound of Preparation 167 (370 mg) is dissolved in 35 mL of ethyl acetate and 6 ml of methanol. To that solution is added 200 mg of 10% Pd on Carbon catalyst and the reaction is hydrogenated under 50 psi of hydrogen for 2 hours. The reaction is evaporated and chromatographed over 15 60 g of silica gel to yield 244 mg of the title compound.

Physical characteristics are as follows:

IR (mineral oil) 3025, 2954, 2871, 2854, 1635, 1619, 1604, 1494, 1456, 1383, 1378, 1256 cm<sup>-</sup>1.

[ $\alpha$ ]<sub>D</sub> (16.764 mg/mL in CH<sub>3</sub>OH)=+39°. Mass spectrum: molecular ion at 393. Anal. found: C, 75.79; H, 8.05; N, 3.27.

## **EXAMPLE 378**

(3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide (Formula X-13 where  $R_1$  is phenethyl) Refer to Chart X

The product of Preparation 168 (156 mg) is added to 5 mL of methylene chloride. To that solution is added 96  $\mu$ L of pyridine and then the reaction is cooled to 0° C. To the aforementioned solution is added 102 mg of 5-trifluoromethyl-2-pyridinyl sulfonyl chloride. The reaction is stirred for 1 hour and then poured into ethyl acetate, washed with water, brine and dried with MgSO<sub>4</sub>. The solvent is evaporated in vacuo and the resulting material chromatographed over 100 g of silica gel eluting with 50% ethyl acetate/hexane to yield 200 mg of the title compound.

Physical characteristics are as follows:

Mass spectrum: molecular ion at 602.

IR (mineral oil) 2953, 2922, 2870, 2853, 1642, 1605, 1459, 1457, 1326, 1259, 1180, 1171, 1141 cm<sup>-</sup>1.

UV (EtOH)  $\lambda_{max}$  ( $\epsilon$ ) 216 (22300), 264 sh (10700), 270 (11500), 279 (12100)

Anal. found: C, 57.53; H, 5.98; N, 4.84.

## **EXAMPLE 379**

 $(3R,6S)-N-[3-[1-(5 6-Dihydro-4-hydroxy-2-oxo-6-propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide (Formula X-13 where <math>R_1$  is phenethyl) Refer to Chart X

The product of Example 378 is added to isopropanol and injected onto a 0.46×25 cm Cyclobond I 2000 column (Advanced Separations Technologies, Inc., Whippany, N.J.). The column is in an ice-water bath. The sample is eluted at 1.0 mL/min. with acetonitrile containing 0.1% diethylamine 55 and 0.6% glacial acetic acid (V/V). The monitor is set at 250 nm. The earlier eluting diastereomer is identical to the compound of Example 298. The second eluting diastereomer is purified over 60 g of silica gel eluting with 40% ethyl acetate/hexane to afford 13 mg of the title compound. 60

Physical characteristics are as follows:

Opposite stereochemistry at C-6 to the compound of Example 298.

<sup>1</sup>H-NMR (CD<sub>3</sub>OD, δ) 8.91, 8.19, 8.16, 8.02, 7.99, 7.25, 7.18, 7.15, 7.13, 7.11, 7.04, 6.97, 6.89, 6.75, 3.95, 2.69, 2.64, 65 2.53, 2.48, 2.13, 1.91, 1.71, 1.68, 1.37, 1.19, 1.17, 1.14, 0.94, 0.92, 0.89, 0.85, 0.83, 0.80, 0.93.

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#### PREPARATION 169

(3S)-3-[(3-Bis(phenylmethyl)amino)phenyl]-4,4-dimethylpentanoic acid methyl ester (Formula LLL-9) Refer to Chart LLL

To anhydrous methanol (2 mL) at room temperature is added titanium (IV) chloride (0.07 mL). The resulting light green solution is stirred for 2 h, treated with the compound of formula LLL-2 wherein R is phenyl (100 mg), prepared by procedures analogous to those described in Chart FF, and refluxed for 18 h. The reaction mixture is allowed to cool and is partitioned between 1N HCl and diethyl ether. The organic layer is separated washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo. Purification by flash chromatography eluting with hexane/ethyl acetate (95:5) affords the title compound (58 mg) as a light amber oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.32–7.20, 7.04, 6.61–6.48, 4.61, 3.48, 2.85–2.80, 2.72–2.55, 0.75 ppm

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ173.69, 148.45, 142.51, 138.91, 128.53, 128.17, 126.78, 117.98, 114.49, 110.89, 54.54, 52.24, 51.40, 35.56, 33.65, 27.87 ppm MS (EI) m/z 415.

#### PREPARATION 170

(3S)-3-[(3-Bis(phenylmethyl)amino)phenyl]-4,4-dimethylpentanoic acid (Formula LLL-10) Refer to Chart LLL

The compound of formula LLL-9 (406 mg) of Preparation 169 is slurried in glacial acetic acid (2.6 mL) and 6N sulfuric acid. The reaction mixture is refluxed for 5 h, allowed to cool and is partitioned between water and diethyl ether. The aqueous layer is separated and extracted two more times with diethyl ether. The combined organic layers are washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo. The resulting light brown residue is dissolved in diethyl ether and treated with dicyclohexylamine (0.16 mL) at 0° C. The solids are isolated, washed with diethyl ether and dried in vacuo. The light brown solid is suspended in diethyl ether and washed with 0.25N HCl. The organic layer is washed with brine, dried over anhydrous sodium sulfate, and concentrated in vacuo, affording the title product (54 mg) as a light brown amorphous solid.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.31–7.19, 7.04, 6.61–6.48, 4.61, 2.81–2.56, 0.74 ppm

<sup>13</sup>C NMR (CDCl<sub>3</sub>) δ179.15, 148.56, 142.28, 138.83, 128.55, 128.23, 126.76, 117.90, 114.49, 110.98, 54.51, 51.83, 35.45, 33.67, 27.84 ppm MS (EI) m/z 401.

## PREPARATION 171

N-[(S)-4-Benzyl-2-oxazolidinone]3-aminocinnamate amide (Formula HHH-4) Refer to Chart HHH

A 1 liter round-bottomed flask with nitrogen inlet and addition funnel is charged with 10.02 g of commercially available (S)-4-benzyl-2-oxazolidinone and 260 mL of tetrahydrofuran and then cooled to -78° C. To the aforementioned solution is added 37 mL of n-butyl lithium during which time a white solid separates from the reaction solution. To that suspension is added 11.46 g of trans-3-nitrocinnamic acid chloride (prepared from the treatment of commercially available 3-nitrocinnamic acid with oxalyl chloride) in a small volume of THF. The resulting pale yellow homogeneous solution is allowed to warm to room temperature and quenched with a saturated ammonium chloride solution and is extracted with ethyl acetate. The organic layer is separated, washed with brine and water,

dried over magnesium sulfate, filtered and concentrated to give a reddish brown syrup (formula HHH-3 in Chart HHH) which is used without further purification. The aforementioned crude reaction mixture is added to ethanol containing 64.18 grams of SnCl<sub>2</sub>.2H<sub>2</sub>O and that mixture heated at reflux for 20 minutes. The reaction is cooled to room temperature and poured into ice. The mixture is brought to pH 9–10 with saturated aqueous Na<sub>2</sub>CO<sub>3</sub>. The mixture is filtered and the filter cake washed extensively with ethyl acetate. The filtrate is washed with brine and the organic 10 phase is dried (Na<sub>2</sub>SO<sub>4</sub>), filtered, and concentrated in vacuo to give a yellow solid. Recrystallization from ethanol gives 11.56 g of the title product.

Physical characteristics are as follows:

IR (mineral oil) 3450, 3369, 2924, 1771, 1678, 1620, 15 1462, 1392, 1357, 1347, 1214 cm<sup>-</sup>1

 $[\alpha]_D$  (14.418 mg/mL in CHCl<sub>3</sub>)=+51°

### PREPARATION 172

N-[(S)-4-Benzyl-2-oxazolidinone]3-(bis(phenylmethyl) amino) cinnamate amide (Formula HHH-5) Refer to Chart HHH

The amine of formula HHH-4 from Preparation 172 (10.13 g), 10.48 g of potassium carbonate, 8.3 mL of benzyl bromide and 100 mL of acetonitrile is heated at reflux for 3 hours. The reaction is cooled to room temperature and partitioned between water and ethyl acetate. The aqueous is extracted several additional times with ethyl acetate. The combined ethyl acetate extracts are dried (Na<sub>2</sub>SO<sub>4</sub>), filtered and concentrated in vacuo. The residue is purified via silica gel chromatography eluting with 25% ethyl acetate/hexane to yield 8.87 g of the title product.

Physical characteristics are as follows:

<sup>13</sup>C-NMR (CDCl<sub>3</sub>, ppm) 165, 153, 149, 147, 138, 135, 129.6, 129.3 128.8, 128.6, 127, 126.9, 126.5, 116.54, 116.50, 114, 113, 65, 55, 54, 37

IR (mineral oil) 2954, 2870, 2854, 1776, 1677, 1616, 1595, 1493, 1454, 1353, 1209, 988 cm<sup>-1</sup>

## PREPARATION 173

(3S)(4S) 3-[3-(3-(bis(phenylmethyl)aminophenyl) 40 pentanoyl]-4-phenyl-2-oxazolidinone (Formula HHH-6) Refer to Chart W

A 100-mL, three-necked flask equipped with a stir-bar, 25-mL pressure-equalizing addition funnel, and a nitrogen inlet is charged with copper(I) bromide dimethyl sulfide 45 complex (1.69 g), 20 mL of tetrahydrofuran and 10 mL of dimethyl sulfide. The addition funnel is charged with the title compound of Preparation 172 (2.747 g) and 10 mL of tetrahydrofuran. The reaction mixture is cooled to -40° C. and ethyl magnesium bromide (5.5 mL of a 3.0M solution in 50 ether) is added dropwise over 5 min. The resulting black mixture is stirred another 10 min at -40° C. and then allowed to warm to -10° C. The solution of the title compound of Preparation 172 in tetrahydrofuran is added dropwise to the reaction mixture over 17 min. The addition funnel is then 55 rinsed with another 3 mL of tetrahydrofuran, and the reaction mixture is stirred for 2.5 h at ca. -40° to -60° C. The reaction is quenched by pouring the mixture into 50 mL of saturated aqueous ammonium chloride solution, and the organic solvents are removed by concentration in vacuo. The 60 resulting residue is partitioned between 75 mL of ethyl acetate and 50 mL of water and filtered through glass wool. The organic layer is then separated, washed with two 100-mL portions of 10% ammonium hydroxide solution and 50 mL of brine, dried over magnesium sulfate, filtered and 65 concentration in vacuo to yield 3.59 g of a yellow oil. Column chromatography on 150 g of silica gel (elution with

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5-15% ethyl acetate/hexane) affords two diastereomeric products. 1.602 g of the title compound (the less polar diastereomer) is isolated as a pale yellow oil

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>) 87.32–7.17, 7.06, 6.60, 6.55, 4.63, 4.43–4.37, 4.00, 3.85, 3.37, 3.20, 3.08, 3.02–2.92, 2.62, 1.71–1.48, 0.73 ppm]

Also isolated from the column is 0.310 g of the more polar diastereomer as a pale yellow oil.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CDCl<sub>3</sub>)  $\delta$ 7.32–7.18, 7.12, 7.05, 6.64–6.56, 4.63, 4.60–4.52, 4.08–4.04, 3.48–3.38, 3.07–2.96, 2.48, 1.69–1.48, 0.73 ppm.

In addition, fractions containing 0.708 g of a ca. 1:4 ratio mixture of the less polar to more polar diastereomers are collected from the column.

## PREPARATION 174

(3S,6S)-3-[1-(3-aminophenyl)propyl]-5,6-dihydro-4hydroxy-6-(2-phenylethyl)-6-propyl-2H-pyran-2-one (Formula FFF-6) Refer to Chart FFF

Following the general procedure of Preparation 162, and making non-critical variations, but substituting the title product of Preparation [U-141164] for the title product of Preparation 161, 0.040 g of crude title compound is obtained. This compound is used immediately in the next step without further purification.

## **EXAMPLE 380**

(3S,6S)-N-[3-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide (Formula FFF-7) Refer to Chart FFF

Following the general procedure of Example 374, and making non-critical variations, but substituting the title product of Preparation 174 for the title product of Preparation 162, 0.015 g of the title compound is obtained as a white foam.

Physical characteristics are as follows:

<sup>1</sup>H NMR (CD<sub>3</sub>OD) δ8.95, 8.25–8.21, 8.07–8.02, 7.25–6.93, 3.94–3.88, 2.70–2.51, 2.20–2.18, 1.97–1.66, 1.40–1.30, 0.92–0.81 ppm.

Thus, for example, the compounds of the present invention include the following individual stereoisomers:

- 5-cyano-N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R)-(3,3,3-trifluoropropyl)-2H-pyran-3-y1]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- 5-cyano-N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- 5-cyano-N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(S)-(3,3,2-trifluoropropyl)-2H-pyran-3-y1]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- 5-cyano-N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2, 2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2, 2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,

- N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(S)-(2-phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2, 2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide.
- N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(R)-(2-5 phenethyl)-6-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2, 2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- 5-amino-N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2pyridinesulfonamide,
- 5-amino-N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- 5-amino-N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- 5-amino-N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(S)-(3,3,3-trifluoropropyl)-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R)-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(S)-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(R)-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]phenyl]-2-pyridinesulfonamide,
- N-[3-(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenethyl)-6-(S)-propyl-2H-pyran-3-yl]-2,2-35 dimethylpropylphenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(R)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3-(R)-[-[4-hydroxy-2-oxo-6,6-din-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-din-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(R)-[1-[4-hydroxy-2-oxo-6,6-din-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]1-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-din-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2dimethylpropyl]-phenyl]1-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(R)-[1-[4-hydroxy-2-oxo-6,6-di-phenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-diphenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3-(R)-[1-[4-hydroxy-2-oxo-6,6-di-phenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-di-65 phenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,

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- 4-Trifluoromethyl-N-[3-(R)-[1-[4-hydroxy-2-oxo-6,6-din-propyl-5,6-dihydro-2H-pyran-3-yl]-2,2dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-di-hydro-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 4-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(R)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(R)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(S) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- N-[3(R)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(R)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3(R)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(S)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3(S)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(R)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,

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- N-[3(S)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(S)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- 5-Amino-N-[3(R)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(S) 5 -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(S) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6 (R)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6 (R)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl) phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(S) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl) phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R) -(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl) 25 phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(S) -(2-phenylethyl)-6-propyl- 2H-pyran-3-yl]propyl) phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R)-(1-[6(R)-(2-[4-fluorophenyl]ethyl)-5, 6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(R)-(1-[6(S)-(2-[4-fluorophenyl]ethyl)-5, 6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[6(R)-(2-[4-fluorophenyl]ethyl)-5, 6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[6(S)-(2-[4-fluorophenyl]ethyl)-5, 40 6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide,
- N-[3(R)-(1-[5,6-Dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3(S)-(1-[5,6-Dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- 5-Amino-N-[3(R)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) 50 phenyl]-2-pyridinesulfonamide,
- 5-Amino-N-[3(S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(R)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) phenyl]-2-pyridinesulfonamide,
- 5-Cyano-N-[3(S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl) 60 phenyl]-2-pyridinesulfonamide,
- N-[3(R)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3(S)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-65 hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-1methyl-1H-imidazole-4-sulfonamide,

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- N-[3(R)-(1-[6,6-Bis(2-phenyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide,
- N-[3(S)-(1-[6,6-Bis(2-phenyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide.
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide.
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-aminopyridine-2-sulfonamide,

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-aminopyridine-2-sulfonamide,

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-5phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide.

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl) - 2 H - pyran-3-yl) - 2,2-dimethylpropyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,

N- $\{3-\{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl\}-2,2-dimethylpropyl\}$ phenyl $\}$ -5-cyanopyridine-2-sulfonamide.

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)-2,2-dimethylpropyl}phenyl]-5-cyanopyridine-2-sulfonamide

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-sulfonamide.

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-25 sulfonamide

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-aminopyridine-2-sulfonamide.

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-30 2H-pyran-3-yl)-propyl}phenyl]-5-aminopyridine-2-sulfonamide.

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide,

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide,

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) 40 propyl}phenyl}-5-cyanopyridine-2-sulfonamide,

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-cyanopyridine-2-sulfonamide,

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-1-methyl-1H-imidazole-4-sulfonamide,

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-1-methyl-1H-imidazole-4-60 sulfonamide,

N-[3-{1(R)-(4-Hydroxy-5,6-dibydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-aminopyridine-2-sulfonamide,

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-65 fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-aminopyridine-2-sulfonamide,

N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-aminopyridine-2-sulfonamide,

N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-(2-(4-fluorophenyl)ethyl)-6-propyl-2H-pyran-3-yl) propyl}phenyl]-5-aminopyridine-2-sulfonamide,

$$R_1$$
 $R_2$ 
 $R_3$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_3$ 

$$R_1$$
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_4$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_5$ 
 $R_7$ 

$$R_1$$
 $R_2$ 
 $R_3$ 
 $R_2$ 
 $R_0$ 
 $R_3$ 

$$R_{10}$$
  $R_{3}$   $R_{20}$   $O$ 

$$\begin{array}{c|c} & & & \text{IIA} \\ \hline R_{10} & & & \\ \hline R_{20} & & & \\ \hline O & & & \\ \hline \end{array}$$

$$\begin{array}{c} O \\ R_{10} \\ \hline \\ R_{20} \\ O \end{array} \qquad \begin{array}{c} O \\ O \\ \end{array}$$

$$(CH_2)_p$$

$$\begin{matrix} OH & VI \\ R_2 & \\ R_6 & O \end{matrix}$$

179
-continued

J

180

$$(CH_2)_n \longrightarrow O$$

$$O$$

$$O$$

$$O$$

$$O$$

## CHART B

185

-continued

CHART C

186 -continued CHART D

D-5

CHART E

$$(R \text{ or } S)$$

$$HO$$

$$N$$

$$N$$

$$N$$

$$N$$

$$N$$

$$E-10$$

CHART F

-continued

10

15

20

-continued CHART F -continued CHART F

#### CHART J

-continued CHART J

-continued CHART M

-continued CHART N

60

$$R-1$$
 $NO_2$ 
 $R-2$ 
 $NH_2$ 

CHART S

-continued
CHART S

## CHART V

$$\begin{array}{c}
\downarrow \\
\downarrow \\
\downarrow \\
\downarrow \\
V-1
\end{array}$$

$$\begin{array}{c}
\downarrow \\
\downarrow \\
\downarrow \\
V-2
\end{array}$$

$$H_3$$
C  $W$ -9

$$H_3C$$
 $OH$ 
 $O$ 
 $CH_3$ 
 $N(Bn)_2$ 
 $W-10$ 

$$H_3C$$
 $OH$ 
 $OH$ 
 $OH$ 
 $NH_2$ 
 $W-12$ 

### CHART X

-continued CHART X

$$O \bigvee_{\text{N}} \bigcap_{\text{N} \in \mathbb{N}} N(Bn)_2 \quad X \cdot 6$$

$$X_{A}$$
 $O$ 
 $CH_{3}$ 
 $N(Bn)_{2}$ 
 $X-8$ 

$$H_3$$
C $H_3$  $N(Bn)_2$   $X-9$ 

$$R_1$$
OH
O
 $X_A$ 
 $N(Bn)_2$ 
 $X_{-10}$ 

$$x_{A}=0$$
 $N$ 

$$H_3C$$
 $OH$ 
 $O$ 
 $CH_3$ 
 $N(Bn)_2$ 
 $X-10$ 

$$\begin{array}{c|c} & & & \\ & & \\ & & & \\ & & \\ & & \\ & & & \\ & & \\ & & & \\ & & \\ & & \\ & & & \\ & & \\ & & & \\ & &$$

$$H_3C$$
 $OH$ 
 $OH$ 
 $N(Bn)_2$ 
 $X-11$ 

$$H_3C$$
 $OH$ 
 $OH$ 
 $OH$ 
 $NH_2$ 
 $X-12$ 

-continued CHART X

-continued CHART Y

CHARTY

N(Bn)<sub>2</sub>

NH<sub>2</sub>

Y-10

$$O$$

N(Bn)<sub>2</sub>
 $O$ 

N(Bn)<sub>2</sub>
 $O$ 

NH<sub>2</sub>

Y-10

-continued CHART Z

$$\bigvee_{N(Bn)_2}^{CHO} Z-4$$

CHART AA

-continued CHART AA

-continued CHART AA

-continued CHART AA

BB-10

BB-12

-continued CHART BB

-continued CHART BB

CHART CC

CHART CC

CH<sub>3</sub>

$$CH_3$$
 $CC_2(Y-6)$ 
 $CH_3$ 
 $CC_2(Y-6)$ 
 $CC_3$ 
 $CC_3$ 
 $CC_4$ 
 $CC_4$ 
 $CC_4$ 
 $CC_5$ 
 $CC_6$ 
 $CC_7$ 
 $C$ 

CC-5

OTMS 
$$N \equiv C$$
  $N(Bn)_2$   $CC-5$ 

$$H_3$$
C  $OH$   $N(Bn)_2$   $CC-8$ 

### CHART DD

DD-5

-continued CHART DD

OTMS 
$$CH_3$$

$$N \equiv C \qquad \qquad N(Bn)_2 \qquad DD-5$$

$$N(Bn)_2$$
 DD-8

$$H_3C$$
OH
 $OH$ 
 $N(Bn)_2$ 
 $DD-10$ 

### -continued CHART DD

### CHART EE

CHART EE

CH3

CH3

N(Ba)<sub>2</sub>

EE-3

$$(CH_3)$$

N(Ba)<sub>2</sub>

EE-4

N(Ba)<sub>2</sub>
 $(CH_3)$ 

N(Ba)<sub>2</sub>

EE-5

-continued CHART EE

$$N \equiv C \begin{tabular}{ll} CH_3 \\ \hline N \equiv C \begin{tabular}{ll} N(B_n)_2 \\ \hline EE-5 \end{tabular}$$

$$N(Bn)_2$$
 EE-8

# -continued CHART EE

# CHART FF

-continued CHART FF

# -continued CHART GG

-continued CHART GG

$$H_3C$$
 $CH_3$ 
 $CH_3$ 
 $OH$ 
 $NH_2$ 
 $GG-15$ 
 $CH_3$ 
 $C$ 

CHART HH

НН-3

25 -continued CHART HH N(CH<sub>2</sub>Ph)<sub>2</sub> 30 OTs HH-4 35 N(CH<sub>2</sub>Ph)<sub>2</sub> SPh HH-5 45 HH-6 (BB-7) N(CH<sub>2</sub>Ph)<sub>2</sub> 55 SPh OH 60 HH-7

65

-continued CHART HH

HH-8

$$\stackrel{\text{OH}}{\longrightarrow} \stackrel{\text{NH}_2}{\longrightarrow}$$

HH-9

HH-10

CHART JJ

N(CH<sub>2</sub>Ph)<sub>2</sub> JJ-1 (X-6) N(CH<sub>2</sub>Ph)<sub>2</sub> JJ-2 N(CH<sub>2</sub>Ph)<sub>2</sub> OH 31-3 N(CH<sub>2</sub>Ph)<sub>2</sub> OTs JJ-4 N(CH<sub>2</sub>Ph)<sub>2</sub> SPh JJ-5 JJ-6 (BB-7)

-continued CHART JJ N(CH<sub>2</sub>Ph)<sub>2</sub> 10 JJ-7 15 N(CH<sub>2</sub>Ph)<sub>2</sub> 20 25 JJ-8 30 N(CH<sub>2</sub>Ph)<sub>2</sub> J**J-**9 40 45 50 JJ-10 55 60

JJ-11

-continued CHART KK

KK-4

$$\bigcup_{OH} \bigcup_{O} \bigvee_{NH_2} \longrightarrow$$

KK-6

KK-5

5
$$OH \qquad H \\ N-SO_2 \qquad N$$

$$KK-7$$

CHART LL

LL-2

LL-9

-continued CHART LL

5
$$OH \longrightarrow H \\ N-SO_2 \longrightarrow N$$

$$ILL-11$$

15 CHART MM

$$\begin{array}{c} \text{SPh} \\ \text{50} \end{array}$$

65

-continued CHART MM

OH N(CH<sub>2</sub>Ph)<sub>2</sub>

MM-5

OH

NH<sub>2</sub>

NH<sub>2</sub>

O NN-1 (GG-5)

NN-2

-continued CHART NN

5 R O N(CH<sub>2</sub>Ph)<sub>2</sub>
OH
NN-3

15 R O N(CH<sub>2</sub>Ph)<sub>2</sub>
OTs
20 NN-4

25 R O N(CH<sub>2</sub>Ph)<sub>2</sub>
30 SPh NN-5

35 NN-6 (BB-7)

45 N(CH<sub>2</sub>Ph)<sub>2</sub>

NN-7

SPh N(CH<sub>2</sub>Ph)<sub>2</sub>

60 NN-8

-continued CHART NN

$$\stackrel{\text{OH}}{\longrightarrow} \stackrel{\text{OH}}{\longrightarrow} \stackrel{\text{NH}_2}{\longrightarrow}$$

NN-9

$$\bigcap_{O} \bigcap_{N \to O_2} \bigcap_{N} \bigcap_{N \to O_2} \bigcap_{N$$

NN-10

00-3

00-4

00-5

XX-3 (BB-7)

## CHART YY

$$\begin{array}{c|c}
R & O & N(CH_2Ph)_2 \\
\hline
NMe_2 & ZZ-2
\end{array}$$

## CHART AAA

AAA-5 (KK-5)

BBB-6 (LL-9)

-continued CHART DDD

CHART EEE

CHART GGG

-continued CHART GGG

-continued CHART GGG

-continued CHART GGG

# CHART HHH

-continued CHART HHH

-continued CHART HHH

-continued CHART HHH

$$CH_3$$
 $CH_3$ 
 $NHSO_2$ 
 $NHSO_2$ 
 $NHSO_2$ 
 $NHH-13$ 

HHH-12

CHART III

-continued CHART III

-continued CHART III

$$N(Bn)_2$$
 III-8

 $N(Bn)_2$  III-9

 $N(Bn)_2$  III-10

 $N(Bn)_2$  III-10

 $N(Bn)_2$  III-11

 $N(Bn)_2$  III-11

 $N(Bn)_2$  III-12

 $N(Bn)_2$  III-12

 $N(Bn)_2$  III-13

-continued CHART III

CHART JJJ

JJJ-3

O NLi

O NLi

O CH<sub>3</sub>

JJJ-4

-continued CHART JJJ

$$\begin{array}{c|c} O & O & CH_3 \\ \hline O & N & N(Bn)_2 \\ \hline \end{array}$$
 JJJ-6

-continued CHART JJJ

323

CHART KKK

JJJ-14 (From JJJ-5b)

-continued CHART KKK

$$R_{1} \longrightarrow CH_{5} \longrightarrow KKK-2$$

$$R_{1} \longrightarrow CH_{5} \longrightarrow KKK-3$$

$$R_{1} \longrightarrow KKK-4b \longrightarrow KKK-4b$$

$$R_{2} \longrightarrow KKK-4b$$

$$R_{3} \longrightarrow KKK-4b$$

$$R_{4} \longrightarrow KKK-4b$$

$$R_{5} \longrightarrow KKK-5$$

$$R_{1} \longrightarrow KKK-5$$

$$R_{1} \longrightarrow KKK-5$$

$$R_{2} \longrightarrow KKK-5$$

$$R_{3} \longrightarrow KKK-7b$$

$$R_{4} \longrightarrow KKK-7b$$

$$R_{5} \longrightarrow KKK-7b$$

$$R_{5} \longrightarrow KKK-7b$$

$$R_{6} \longrightarrow KKK-7b$$

$$KKK-7b \longrightarrow KKK-7b$$

$$KKK-7b \longrightarrow KKK-7b$$

$$KKK-7b \longrightarrow KKK-7b$$

-continued CHART KKK

$$X_{a} = 0$$

$$X_{b} = 0$$

$$X_{b$$

-continued CHART KKK

$$R_1$$
 $R_2$ 
 $OH$ 
 $OH$ 
 $NHSO_2R_4$ 
 $KKK-14b$ 

$$X_{B} = 0$$
 $N$ 
 $N$ 

$$R_1$$
OH
O
 $X_B$ 
 $N(Bn)_2$ 
 $KKK-16$ 

KKK-15(X-6)

-continued
CHART KKK

$$R_{1} \longrightarrow CH_{3} \longrightarrow KKK.17$$

$$R_{1} \longrightarrow CH_{3} \longrightarrow KKK.18$$

$$KKK.18a \longrightarrow KKK.18a$$

$$KKK.18b \longrightarrow KKK.18a$$

$$KKK.18a \longrightarrow KKK.18a$$

-continued CHART KKK

$$X_{g} = O$$

$$N$$

$$N_{g} = O$$

### -continued CHART KKK

$$R_1$$
 $R_2$ 
 $OH$ 
 $NH_2$ 
 $KKK-13$ 
 $OH$ 
 $CH_3$ 
 $OH$ 
 $NHSO_2R_4$ 
 $KKK-14a$ 

### CHART LLL

-continued CHART LLL

$$H_{j,C} \xrightarrow{CH_{j}} \xrightarrow{N(Ba)_{2}} \xrightarrow{LLL-18} H_{j,C} \xrightarrow{N(Ba)_{2}} \xrightarrow{LLL-18} H_{j,C} \xrightarrow{N(Ba)_{2}} \xrightarrow{LLL-18} H_{j,C} \xrightarrow{N(Ba)_{2}} \xrightarrow{LLL-18} H_{j,C} \xrightarrow{N(Ba)_{2}} \xrightarrow{N(Ba)_{2}} \xrightarrow{LLL-18} H_{j,C} \xrightarrow{N(Ba)_{2}} \xrightarrow{N(Ba)_{2}}$$

-continued

-continued CHART LLL

# CHART MMM

### MMM-1 (X-11)

$$\begin{array}{c} \text{OH} \\ \text{CH}_3 \\ \text{N(CH}_2\text{Ph})_2 \\ \text{AND} \end{array}$$

### MMM-2

# OH CH<sub>3</sub> NH<sub>2</sub>

### MMM-4

### MMM-5

CH<sub>3</sub>

### MMM-6

MMM-7

H<sub>3</sub>C

MMM-3

он

347 CHART NNN NNN-1 (W-2) NHMe нο NNN-3

NNN-4

но

NNN-10 (W-13)

60

CHART OOO

CHART PPP

-continued CHART PPP

CHART QQQ

-continued CHART QQQ

-continued CHART QQQ

$$\begin{array}{c} \text{Ph} \\ \text{Ph} \\ \text{CF}_3 \end{array} \begin{array}{c} \text{R}_1 \\ \text{H} \\ \text{N}_{SO_2} \end{array} \begin{array}{c} \text{Ph} \\ \text{CF}_3 \\ \text{QQQ-16} \end{array}$$

CHART RRR

$$P_{h}$$

$$CH_{3}$$

$$RRR-A$$

$$RRR-B$$

$$RRR$$

-continued CHART RRR

-continued CHART RRR

### CHART SSS

SSS-A SSS-B

$$H_3C$$
 $O$ 
 $O$ 
 $NH$ 
 $R_2-SO_2CI$ 

-continued CHART SSS

$$H_3C$$
 $CH_3$ 
 $CSS-7$ 

$$R_2$$

ТТТ-А

TTT-4

 $R_3$ -SO<sub>2</sub>CI

$$R_2$$
 $R_2$ 
 $R_2$ 
 $R_2$ 
 $R_2$ 
 $R_3$ 
 $R_4$ 
 $R_5$ 
 $R_5$ 
 $R_7$ 
 $R_7$ 

CHART UUU

-continued CHART TTT

$$\begin{array}{c|c} & \text{HO} & R_1 \\ \hline \\ R_2 & & \\ \hline \\ \text{TTT-6} & \\ \end{array}$$

50

55

60

65

UUU-5

OН

HN ОН D-5

CHART WWW

WWW-1

-continued CHART WWW

-continued CHART XXX www-2

OH 
$$R_3$$

OH  $R_3$ 

ОН

-continued CHART YYY -continued CHART ZZZ

$$\begin{array}{c} OH \\ R_1 \\ R_2 \end{array} \begin{array}{c} OH \\ OH \\ R_1 \\ R_2 \end{array} \begin{array}{c} OH \\ OH \\ R_1 \\ R_2 \end{array} \begin{array}{c} OH \\ OH \\ R_1 \\ R_2 \end{array} \begin{array}{c} OH \\ OH \\ R_1 \\ R_2 \end{array} \begin{array}{c} OH \\ OH \\ R_3 \end{array}$$

CHART ZZZ

$$SYY-2$$
 5  $Br$   $TMS$   $ZZZ-3$ 

ZZZ-1

10

-continued CHART AAAA

AAAA-9 R O O NBn<sub>2</sub>

$$CISO_2$$
 $CISO_2$ 
 $CF_3$ 
 $CF_3$ 

BBBB-4 R OH NB
$$_{n_2}$$
 CI-SO<sub>2</sub> CF<sub>3</sub> BBBB-5

-continued
CHART BBBB

OH

NHSO<sub>2</sub>

N =

BBBB-6

15 CHART CCCC 20 NBn<sub>2</sub> 25 30 NBn<sub>2</sub> 35 он CCCC-3 40 NBn<sub>2</sub> 45 50 CCCC-4 NBn<sub>2</sub> 55

15

-continued CHART CCCC -continued

OH 
$$R_1$$
 DDDD-5

DDDD-6

20 OH 
$$R_1$$
 DDDD-7  $CH_2$   $OH_2$   $OH_3$   $OH_4$   $OH_5$   $OH$ 

CHART DDDD

$$(CH_2)_n$$
  $O$   $NO_2$   $OH$   $R_1$   $NO_2$   $NO_2$ 

DDDD-3 
$$H_3C$$
  $OH$   $CH_3$  EEEE-2  $C \equiv N$ 

-continued CHART EEEE

$T\Delta$	RI	F	I-continued

	CHARI	EEEE				IADLE 1-	СОППППЕС	
	OH CI	Н3	EEEE-4		Compound of	н	V Protease FITC Ass	ay
	OH	^		5	Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)
H <sub>3</sub> C					98 Second Compound			0.800 0.840
	0 0	Y			Develor Consposito			0.800
		ŃН 		10	139	0.123	98.33	
$\searrow$ $^{\prime\prime}$		l				0.370	101.22	
<b>&gt;</b>		_ SO₂				1.100	104.71	
						3.300	99.3	
	N N					10.000 30.000	99.28 102.85	
F <sub>3</sub> C				1.5		30.000	102.83	1.890
				15	140	0.123	103.22	1.000
OН	CH₃		EEEE-5			0.370	96.01	
						1.100	107.37	
H <sub>3</sub> C	$\sim$	5				3.800	112.51	
\						10.000	112.53	
		ال		20		30.000	119.14	4 440
	*° <b>*</b>				40	0.122	59.6	1.440
H <sub>3</sub> C	NH				40	0.123 0.370	101.71	
	O <sub>2</sub> S					1.100	98.73	
	Ĩ					3.300	105.16	
	,, <i>/</i>					10.000	88.7	
	N			25		30.000	72.74	
	L J							10.800
					41	0.123	103	
	l CF₃					0.370	102.38	
	CF3					1.100	103.92 100.93	
				30		3.300 10.000	85.88	
	TAB	LĖI		30		30.000	72.79	
	11 115							3.170
Compound of	н	V Protease FTTC Ass	av		44	0.123	98.43	
•						0.370	114.5	
Example No.	Dose (uM)	Protease % Inhib	$K_i$ $(nM)$			1.100	119.79	
126	0.103	71.65		35		3.300	112.7	
136	0.123 0.370	71.65 85.67				10.000 30.000	101.66 80.02	
	1.100	99.02				30.000	80.02	1.800
	3.300	100.99			145B	0.123	81.81	2.000
	10.000	102.37				0.370	88.38	
	30.000	101.94		40		1.100	96.54	
			1.320	40		3.300	87.85	
145A	0.123	108.66				10.000	102.12	
	0.370 1.100	111.34 118.54				30.000	84.52	1.240
	3.300	115.43			135	0.123	33.21	1.240
	10.000	113.05			155	0.370	84.5	
	30.000	114.19		45		1.100	99.09	
			1.100			3.300	96.86	
137	0.123	98.83				10.000	101.49	
	0.370	91.54				30.000	102.4	
	1.100	100.7 109.9			104	0.103	-10	0.480
	3.300 10.000	98.17		50	104	0.123 0.370	<10 61.68	
	30.000	93.82		50		1.100	81.78	
	55.550		0.520			3.300	93.28	
			0.700			10.000	96.4	
138	0.123	100.88				30.000	109.22	
	0.370	95.51						1.600
	1.100	101.11		55	48	0.123	111.37	
	3.300 10.000	99.64 94.75				0.370 1.100	103.64 110.44	
	30.000	104.68				3.300	89.27	
	20.000	104.00	0.730			10.000	110.97	
			1.400			30.000	105.44	
97	0.123	104.87		60				0.520
	0.370	106.06		60	49	0.123	111.16	
	1.100	110.44				0.370	119.71	
	3.300	106.67				1.100	120.17	
	10.000	115.76 115.47				3.300	106.02	
	30.000	115.47	1.000			10.000 30.000	108.34 112.5	
98			0.740	65		30.000	114.3	0.960
First Compound			/ <del>-</del>		50	0.123	100.54	5.500
•								

	-continued

TABLE I-continued

	TABLE I-	-continued		_	TABLE I-continued				
Compound of	Н	IV Protease FITC Ass	ay	_	Compound of	н	IV Protease FITC Ass.	ау	
Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)	5	Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)	
	0.370	108.31		_		30.000	115.02		
	1.100	112.66						0.360	
	3.300	112.42			58	0.123	64.87		
	10.000	101.02				0.370	83.71		
	30.000	84.79	1 700	10		1.100	94.24		
105	0.122	101.06	1.780			3.300	95.88		
103	0.123 0.370	101.26 114.56				10.000 30.000	100.27 89.81		
	1.100	107.19				30.000	09.01	3.800	
	3.300	110.88			59	0.123	76.69	5.000	
	10.000	111.16		4.5	3,	0.370	90.54		
	30.000	110.6		15		1.100	101.9		
			0.880			8.300	99.87		
52	0.123	85.08				10.000	105.16		
	0.370	87.32				30.000	102.02		
	1.100	92.64						3.500	
	3.300	97.38		20	60	0.123	73.03		
	10.000	97.15		20		0.370	94.3		
	30.000	88.89				1.100	101.28		
			1.400			3.300	100.84		
53	0.123	88.61				10.000	105.68		
	0.370	97.74				30.000	107.38		
	1.100	97.95		25		0.100	04.00	0.950	
	3.300	99.62		23	61	0.123	86.83		
	10.000 30.000	90.16 84.37				0.3 <b>7</b> 0 1.100	95.51 103.35		
	30.000	04.37	0.900			3.300	103.54		
55	0.123	<10	0.900			10.000	105.61		
55	0.370	18.77				30.000	103.53		
	1.100	58.27		30		20.000	100.00	0.710	
	3.300	86.98		-	93A	0.123	59.48	*****	
	10.000	98.33				0.370	90.42		
	30.000	85.88				1.100	103.54		
			1.700			3.300	108.54		
107	0.123	92.69				10.000	109.19		
	0.370	99.24		35		30.000	96.57		
	1.100	105.15						6.060	
	3.300	103.44			143	0.123	80.78		
	10.000	110.33				0.370	97.65		
	30.000	103.47	0.000			1.100	104.91		
			0.890			3.300	102.39		
99	0.123	85.69	0.700	40		10.000 30.000	101.25 103.08		
77	0.370	101.55				30.000	103.06	0.800	
	1.100	108.05			144	0.123	80.58	0.000	
	3.300	100.05			• • •	0.370	87.39		
	10.000	106.61				1.100	93.82		
	30.000	103.12				3.300	100.01		
			0.660	45		10.000	98.12		
141	0.123	78.72				30.000	95.88		
	0.370	88.65						1.200	
	1.100	92.04			145	0.123	73.63		
	3.300	88.26				0.370	89.78		
	10.000	97.8		50		1.100	99.69		
	30.000	98.48	1 400	50		3.300	94.8		
142	0.123	78.01	1.400			10.000	96.85 87.97		
142	0.123	78.01 92.52				30.000	01.91	0.490	
	1.100	106.64			100	0.123	102.53	0.490	
	3.300	105.15			100	0.123	102.53		
	10.000	110.58		55		1.100	91.01		
	30.000	106.77		33		3.300	96.54		
			1.600			100.000	100.86		
56	0.123	104.11				30.000	100.62		
	0.370	108.31						0.730	
	1.100	105.31			62	0.123	76.18		
	3.300	105.47		60		0.370	85.15		
	10.000	114.94		00		1.100	85.28		
	30.000	111.25				3.300	78.67		
		20.22	0.230			10.000	79.69		
57	0.123	99.07				30.000	79.39		
	0.370	105.17			100	0.400	102.42	0.800	
	1.100	110.68		65	108	0.123	103.43		
	3.300 10.000	97.8 104.74		55		0.370	102.13		
	10.000	104.74				1.100	101.87		

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70 · 70 F	_		
TABL	-	I-conti	nued
IADL	_	I-COHU	шиси

TABLE I-continued

	TABLE I	-continued				TABLE I-	continued.	
Compound of	н	IV Protease FITC Ass	ay		Compound of	Н	V Protease FITC Ass	ау
Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)	5	Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)
	3.300	102.41				0.370	87.3	
	10.000	107.73				1.100	97.1	
	30.000	106.39				3.300	96.76	
			0.160			10.000	99.68	
109	0.123	105.42		10		30.000	97.43	
	0.370	99.35						15.000
	1.100	103.75			101	0.123	70.24	
	3.300	100.96				0.370	83.98	
	10.000	108.56				1.100	93.35	
	30.000	109.31				3.300	97.01	
239	0.123	83.64		15		10.000	102.48	
	0.370	96.63				30.000	97.35	0.660
	1.100	98.41 99.53			146	0.123	68.12	0.000
	3.300	103.21			140	0.123	87.38	
	10.000					1.100	103.18	
	30.000	108.02	1.440			3.300	103.16	
			0.860	20		10.000	102.54	
152	0.123	11.52	0.800			30.000	101.95	
132	0.370	80.2				50.000	101.75	0.690
	1.100	95.79			147	0.123	77.45	0.050
	3.300	94.43			4	0.370	102.86	
	10.000	95.45				1.100	111.6	
	30.000	96.47		25		3.300	110.34	
			0.710			10.000	114.04	
8	0.123	99.23				30.000	108.28	
	0.370	110.11						1.000
	1.100	102.93			110	0.123	77.89	
	3.300	110.02				0.370	82.72	
	10.000	105.11		30		1.100	95.11	
	30.000	101.91				3.300	99.1	
			0.350			10.000	99.22	
9	0.123	99.09				30.000	101.27	
	0.370	103.78						3.260
	1.100	104.9						3.630
	3.300	104.69		35	102	0.123	87.11	
	10.000	107.08				0.370	92.73	
	30.000	107.87				1.100	102.21	
			0.420			3.300	110.44	
10	0.123	102.17				10.000	116.72	
	0.370	111.74				30.000	107.83	0.700
	1.100	115.65		40	102	0.102	45.51	0.700
	3.300	119.47			103	0.123	65.51	
	10.000	128.59				0.370	82.58 96.86	
	30.000	130.05	£ 710			1.100		
151	0.123	111 02	5.710			3.300 10.000	100.29 104.76	
151	0.370	111.03 114.59				30.000		
	1.100	117.62		45		50.000	96.05	1.720
	3.300	117.02			194	0.123	<10	1.720
	10.000	116.34			227	0.370	20.03	
	30.000	114.87				1.100	53.89	
	23.000	,	0.360			3.300	75.23	
153	0.123	81.27				10.000	85.48	
	0.370	91.11		50		30.000	85.18	
	1.100	100.49			195	0.123	60.89	
	3.300	104.76				0.370	85.08	
	10.000	102.76				1.100	90.79	
	30.000	100.71				3.300	90.83	
			1.850			10.000	93.14	
154	0.123	99.8		55		30.000	92.69	
	0.370	98.17						3.700
	1.100	99.52			150	0.123	78.42	
	3.300	97.59				0.370	96.45	
	10.000	103.54				1.100	100.07	
	30.000	99.18				3.300	102.81	
			0.220	60		10.000	106.88	
240	0.123	96.32		50		30.000	109.34	
	0.370	100.98						5.900
	1.100	102.71			. 148	0.123	81.35	
	3.300	101.88				0.370	91.68	
	10.000	104.28				1.100	95.57	
	30.000	107.17		45		3.300	90.04	
_			1.300	65		10.000	99.17	
1.	0.123	75.4				30.000	93.52	

TABLE I-continued

TABL	E l-continue	d
	HIV Protease	FT

		TOMMENTO		_		171000	Сопинаса	
Compound of	н	IV Protease FTTC Ass	ay	 -	Compound of	н	IV Protease FITC Ass	ay
Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)	5	Example No.	Dose (uM)	Protease % Inhib	K <sub>i</sub> (nM)
			4.770	_		0.370	86.56	
			18.100			1.100	93.7	
149	0.123	80.51				3.300	98.88	
	0.370	87.52				10.000	99.03	
	1.100	96.32		10		30.000	106.06	
	3.300	92.86						3.290
	10.000	97.12			54	0.123	46.64	
	30.000	95.99				0.370	72.41	
			3.410			1.100	87.91	
			62.700			3.300	89.11	
94	0.123	75.76		15		10.000	87.77	
	0.370	106.6				30.000	91.99	
	1.100	107.3						13.300
	3.300	104.91			146	10.000	102.54	
	10.000	109.2				30.000	101.95	
	30.000	111.29						0.690
			16.370	20	192	0.123	44.04	
95	0.123	91.2		20		0.370	76.28	
	0.370	102.33				1.100	93.96	
	1.100	105.86				3.300	96.93	
	3.300	112.79				10.000	103.33	
	10.000	110.04				30.000	94.38	
	30.000	112.69						7.200
			5.350	25	193	0.123	18.42	
96	0.123	94.17				0.370	40.3	
	0.370	119.36				1.100	77.74	
	1.100	122.12				3.300	98.1	
	3.300	111				10.000	108.41	
	10.000	111.32				30.000	103.17	
	30.000	109.23		30				35.000
	20.000	205120	5.300		11	0.123	78.93	
42	0.123	86.15	2.200			0.370	95.26	
	0.370	102.71				1.100	100.26	
	1.100	98.26				3.300	95.12	
	3.300	102.4				10.000	99.66	
	10.000	91.43		25		30.000	104.39	
	30.000	76.12		35		201000		1.900
	20.000		3.100		12	0.123	75.65	
43	0.123	85.63	0.100			0.370	87.16	
	0.370	99.01				1.100	91.79	
	1.100	95.68				3.300	91.11	
	3.300	96.68				10.000	94.74	
	10.000	101.58		40		30.000	95.69	
	30.000	85.57						2.150
			3.650		13	0.123	68.94	
45	0.123	82.22				0.370	88.07	
	0.370	94.37				1.100	93.98	
	1.100	101.04				3.300	95.51	
	3.300	103.16		45		10.000	98.61	
	10.000	89.76				30.000	104.2	
	30.000	67.5	•					4.150
			4.780		14	0.123	65.67	
46	0.123	85.86				0.370	87.96	
	0.370	99.19				1.100	96.79	
	1.100	103.31		50		3.300	96.56	
	3.300	97.62				10.000	101.77	
	10.000	91.45				30.000	106.39	
	30.000	74.13						6.880
			2.920		15	0.123	77.63	
47	0.123	66.3				0.370	88.45	
	0.370	86.79		55		1.100	92.44	
	1.100	94.7		33		3.300	94.03	
	3.300	100.95				10.000	95.84	
	10.000	98.68				30.000	99.23	
	30.000	84.45						2.800
	22.000		3.000		63	0.123	68.88	
			2.980			0.370	79.56	
51	0.123	98.71	2.700	60		1.100	88.58	
	0.370	103.68				3.300	87.44	
	1.100	104.78				10.000	83.58	
	3.300	101.27				30.000	78.84	
	10.000	95.07			64	0.123	27.95	
	30.000	79.72			<b>V</b> 1	0.370	50.83	
	50.000	15.12	2.660	65		1.100	75.60	
106	0.123	60.94	2.000			3.300	80.88	
100	0.123	····				3.300	00.00	

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TABLE I-continued

TARLE II-continued

	TABLE I-continued						TABLE II-continued						
Compour	nd of	I	IIV Protease FID	C Assay		•	EXAMPLE		DOSE	PROTEASE % INHIB.	FTTC KI (NM)		
Example	No.	Dose (uM)	Protease % In	hib l	ζ <sub>i</sub> (nM)	5	NUMBER	ENZYME HIV-1	30.000	76.85	KI (IVM)		
		10.000	82.03					HIV1TANDEM			0.100		
		30.000	84.39				293	HIV-1	0.123	53.45			
250					1.2			HIV-1	0.370	77.51			
261					0.87			HIV-1	1.100	94.18			
260					2.0	10		HIV-1	3.300	103.03			
258					4.3			HIV-1	10.000	97.41	4.300		
259 256					2.2 8.3		295	HIV-1	30.000	92.01	0.071		
250 257					9.0		281	HIV1TANDEM HIV1TANDEM			0.002		
246					1.7		201	HIV1TANDEM			0.002		
247					1.2	15	285	HIV1TANDEM			0.015		
254					3.0	15	203	HIV-1	0.123	81.8	0.015		
255					1.6			HIV-1	0.370	95.8			
248					4.7			HIV-1	1.100	99.11			
249					0.75			HIV-1	3.300	109.33			
251		0.123	70.84					HIV-1	10.000	104.61			
		0.370	90.56			20		HIV-1	30.000	86.84			
		1.100	97.68			20	286	HIV1TANDEM			13.300		
		3.300	94.5					HIV-1	0.123	34.76			
		10.000	94.16					HIV-1	0.370	68.74			
		30.000	93.24					HIV-1	1.100	89.29			
		0			1.9			HIV-1	3.300	93.11			
253		0.123	94.03			25		HIV-1	10.000 30.000	108.17			
		0.370	96.84			23	207	HIV-1	30.000	95.31	0.038		
		1.100	97.64				287 283	HIV1TANDEM			0.038		
		3.300	95.93 96.95				20 <i>3</i> 296	HIV1TANDEM HIV1TANDEM			0.004		
		10.000 30.000	98.52				290	HIV1TANDEM			0.042		
252		0.123	69.96				231	HIV1TANDEM			0.026		
232		0.370	85.05			30	289	HIV1TANDEM			0.133		
		1.100	89.69			20	290	HIV1TANDEM			1.880		
		3.300	100.57				298	HIV1TANDEM			0.004		
		10.000	96.21					HIV1TANDEM			0.003		
		30.000	91.38					HIV1TANDEM			0.007		
					1.6		266	HIV1TANDEM			0.033		
262		0.123	91.8			35	272	HIV1TANDEM			3.600		
		0.370	96.6			55	270	HIV1TANDEM			0.024		
		1.100	97.13					HIV-1	0.123	75.26			
		3.300	95.4					HIV-1	0.370	85.62			
		10.000	94.17					HIV-1	1.100	93.45			
		30.000	89.18					HIV-1	3.300	96.62			
263		0.123	98.08			40		HIV-1	10.000	94.57			
		0.370	98.99					HIV-1	30.000	82.67			
		1.100	99.1				273	HIV1TANDEM	0.400	40	2.300		
		3.300	98.08					HIV-1		< 10			
		10.000	96.21					HIV-1	0.370	23.24			
264		30.000	88.19					HIV-1	1.100	75.38 94.63			
264		0.123	67.18 75.01			45		HIV-1 HIV-1	3.300 10.000	95.93			
		0.370 1.100	67.71					HIV-1	30.000	91			
		3.300	57.62				276	HIV1TANDEM	30.000	91	33.000		
		10.000	53.69				210	HIV-1	0.123	16.49	55.000		
		30.000	64.58					HIV-1	0.370	38.95			
		20.000	04.50		3.7			HIV-1	1.100	66.1			
265		0.123	33.23		5.1	50		HIV-1	3.300	90.01			
203		0.123	56.33					HIV-1	10.000	90.97			
		1.100	57.78					HIV-1	30.000	87.6			
		3.300	63.69				278	HIV1TANDEM			0.040		
		10.000	80.29					HIV-1	0.123	76.76			
		30.000	80.29 85.64					HIV-1	0.370	86.99			
		30.000	63.04		1.0	55		HIV-1	1.100	95.6			
					1.0			HIV-1	3.300	96.91			
								HIV-1	10.000	93.32			
							0.00	HIV-1	30.000	86.18	0.025		
		TAT	T TO TI				268	HIV1TANDEM HIV1TANDEM			0.835		
		IAB	LE II				271 299		0.123	83.51	0.051		
EVANOUE			B = 0	OTT A CT		60	299	HIV-1 HIV-1	0.123	104.76			
EXAMPLE	ENTON IP	-		TEASE	FITC			HIV-1	1.100	117.95			
NUMBER	ENZYME	Ţ	OSE % I	NHIB.	KI (NM)			HIV-1	3.300	117.93			
280	HIV-1		0.123 9	5.28				HIV-1	10.000	128.03			
200	HIV-1			4.98				HIV-1	30.000	102.89			
	HIV-1			3.01				HIV1TANDEM	20.500	102.07	0.200		
	HIV-1			6.69		65	300	HIVTANDEM			0.100		
	HIV-1			8.64				HIVTANDEM			0.100		
		_						- •					

TABLE II-continued

TABLE II-continued

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	TABL	E II-contin	nued		_		TABL	E II-contir		
EXAMPLE NUMBER	ENZYME	DOSE	PROTEASE % INHIB.	FITC KI (NM)	- . 5	EXAMPLE NUMBER	ENZYME	DOSE	PROTEASE % INHIB.	FITC KI (NM)
•	HIV-1	0.123	90.61				HIV-1	3.300	97.03	
	HIV-1	0.370	99.05				HIV-1	10.000	100.2	
	HIV-1	1.100	111.45				HIV-1	30.000	106.4	
	HIV-1	3.300	109.19			316	HIV1TANDEM			0.357
	HIV-1	10.000	105.56				HIV-1	0.123	75.49	
	HIV-1	30.000	104.91		10		HIV-1	0.370	85.02	
302	HIVTANDEM			1.870			HIV-1	1.100	100.32	
	HIVTANDEM			3.600			HIV-1	3.300	95.46	
	HIV-1	0.123	38				HIV-1	10.000	99.71	
	HIV-1	0.370	65.57				HIV-1	30.000	87.91	
	HIV-1	1.100	89.51			317	HIV1TANDEM			0.040
	HIV-1	3.300	118.39		15		HIV-1	0.123	87.38	
	HIV-1	10.000	104.49		13		HIV-1	0.370	94.14	
	HIV-1	30.000	92.16				HIV-1	1.100	98.45	
304	HIV-1	0.123	92.01				HIV-1	3.300	95.97	
	HIV-1	0.370	93.28				HIV-1	10.000	101.26	
	HIV-1	1.100	96.47				HIV-1	30.000	108.59	
	HIV-1	3.300	100.47			318	HIV1TANDEM			0.019
	HIV-1	10.000	107.61		20	515	HIV-1	0.123	98.06	0.025
	HIV-1	30.000	79.68				HIV-1	0.370	106.35	
	HIV1TANDEM	30.000	15.00	0.100			HIV-1	1.100	101.88	
	HIV1TANDEM			0.050			HIV-1	3.300	88.73	
305	HIV-1	0.123	99.99	0.050			HIV-1	10.000	94.49	
303								30.000	82.83	
	HIV-1 HIV-1	0.370 1.100	110.76 114.35		25	319	HIV-1 HIV1TANDEM	50.000	34.63	29.500
						319		0.123	10.75	29.300
	HIV-1	3.300	110.88				HIV-1 HIV-1	0.123	32.65	
	HIV-1	10.000	102.01							
	HIV-1	30.000	57.83	0.400			HIV-1	1.100	60.14	
20.5	HIV1TANDEM		74.70	0.400			HIV-1	3.300	75.86	
306	HIV-1	0.123	71.79		20		HIV-1	10.000	93.46	
	HTV-1	0.370	82.71		30	220	HIV-1	30.000	74.48	0.071
	HIV-1	1.100	89.3			320	HIV1TANDEM			0.071
	HIV-1	3.300	97.29				HIV1TANDEM			0.050
	HIV-1	10.000	82.59				HIV1TANDEM			0.075
	HIV-1	30.000	53.43			322	HIV1TANDEM			1.070
	HIV1TANDEM			0.040			HIV1TANDEM			1.290
307	HIV-1	0.123	77.39		35	324	HIV1TANDEM			0.156
	HIV-1	0.370	99.85			326	HIV1TANDEM			0.029
	HIV-1	1.100	107.87			328	HIV-1			22.000
	HIV-1	3.300	93.34				HIV-1	0.123	27.81	
	HIV-1	10.000	83.49				HIV-1	0.370	79.47	
	HIV-1	30.000	69.74				HIV-1	1.100	95.45	
	HIV1TANDEM			0.072	40		HIV-1	3.300	96.77	
308	HIV-1	0.123	75.06				HIV-1	10.000	96.78	
	HIV-1	0.370	108.14				HIV-1	30.000	92.17	
	HIV-1	1.100	95.01			329	HIV-1			12.000
	HIV-1	3.300	108.43				HIV-1	0.123	46.4	
	HIV-1	10.000	110.75				HIV-1	0.370	88.19	
	HIV-1	30.000	96.28		4.5		HIV-1	1.100	96.63	
	HIV1TANDEM			0.074	45		HIV-1	3.300	100.32 .	
310	HIV-1	0.123	16.81				HIV-1	10.000	97.07	
	HIV-1	0.370	50.11				HIV-1	30.000	96.35	_
	HIV-1	1.100	78.69			330	HIV1TANDEM			0.524
	HIV-1	3.300	100.22				HIV-1	0.123	93.74	
	HIV-1	10.000	124.77				HIV-1	0.370	94.32	
	HIV-1	30.000	110.91		50		HIV-1	1.100	93.66	
	HIV1TANDEM			1.500			HIV-1	3.300	85.63	
311	HIV-1	0.123	86.51				HIV-1	10.000	87.9	
	HIV-1	0.370	91.49				HIV-1	30.000	69.82	
	HIV-1	1.100	101.8			331	HIV1TANDEM			0.272
	HIV-1	3.300	96.5				HIV-1	0.123	99.76	
	HIV-1	10.000	93.77		55		HIV-1	0.370	104.06	
	HIV-1	30.000	77.63				HIV-1	1.100	108.51	
	HIV1TANDEM			0.007			HIV-1	3.300	99.3	
312	HIV1TANDEM			0.255			HIV-1	10.000	103.28	
314	HIV1TANDEM			0.700			HIV-1	30.000	93.3	
	HIV-1	0.123	82.92			332	HIV1TANDEM			0.400
	HIV-1	0.370	96.14		60		HIV-1	0.123	81.87	
	HIV-1	1.100	114.86		60		HIV-1	0.370	85.65	
	HIV-1	3.300	100.76				HIV-1	1.100	86.23	
	HIV-1	10.000	88.75				HIV-1	3.300	93.28	
	HIV-1	30.000	73.42				HIV-1	10.000	91.68	
315	HIV1TANDEM			0.029			HIV-1	30.000	95.08	
	HIV-1	0.123	79.95				HIV-1			1.600
	HIV-1	0.370	87.25		65	333	HIV-1	0.123	66.73	
	HIV-1	1.100	88.08			233	HIV-1	0.123	85.07	
	*** *-*	1.100	00.00				*** 4.7	0.570	35.01	

TABLE II-continued

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TABLE II-continued

		AL II COLUM									
EXAMPLE NUMBER	ENZYME	DOSE	PROTEASE % INHIB.	FTTC KL (NM)	. 5	EXAMPLE NUMBER	ENZYME	DOSE		PROTEASE % INHIB.	FTTC KI (NM)
	HIV-1	1.100	85.12			-	HIV-1	1.100		71.4	
	HIV-1	3.300	93.69				HIV-1	3.300		83	
	HIV-1	10.000	89.38				HIV-1	10.000		90.97	
	HIV-1	30.000	77.91				HIV-1	30.000		87.18	
	HIV-1			7.700		348	HIV1TANDEM				1.960
334	HIV1TANDEM			0.450	10		HIV-1	0.123		53.47	
	HIV-1	0.123	93.49				HIV-1	0.370		78.32	
	HIV-1	0.370	90.25				HIV-1	1.100		89.84	
	HTV-1	1.100	94.57				HIV-1	3.300		92.96	
	HIV-1	3.300	102.47				HIV-1	10.000		96.28	
	HIV-1	10.000	97.61				HIV-1	30.000		84.67	
	HIV-1	30.000	96.3		15	349	HIV1TANDEM	0.400			0.111
335	HIV-1	0.123	60.07				HIV-1	0.123		74.5	
	HIV-1	0.370	99.75				HIV-1	0.370		88.21	
	HIV-1	1.100	97.05				HIV-1	1.100		99.92 104.99	
	HIV-1	3.300	92.06 89.77				HIV-1	3.300			
	HIV-1	10.000					HIV-1	10.000 30.000		103.49 98.24	
	HIV-1 HIV1TANDEM	30.000	76.25	0.040	20	351	HIV-1 HIV-1	0.123	<	10	
336	HIV-1	0.123	65.64	0.040		331	HIV-1	0.370	`	10	
330	HIV-1	0.370	112				HIV-1	1.100	`	25.4	
	HIV-1	1.100	89.54				HIV-1	3.300		55.11	
	HIV-1	3.300	88.06				HIV-1	10.000		78.53	
	HIV-1	10.000	77.12				HIV-1	30.000		90.55	
	HIV-1	30.000	62.28		25		HIV-1	20.000		30.55	558.000
	HIV1TANDEM	50,000	02.20	0.032		352	HIV-1	0.123	<	10	
338	HIV-1	0.123	61.74	0.002			HIV-1	0.370	-	25.31	
	HIV-1	0.370	85.32				HIV-1	1.100		47.78	
	HIV-1	1.100	80.46				HIV-1	3.300		74.99	
	HIV-1	3.300	89.62				HIV-1	10.000		85.86	
	HIV-1	10.000	83.53		30		HIV-1	30.000		87.82	
	HIV-1	30.000	62.34				HIV-1				168.000
	HIV1TANDEM			0.100		353	HIV-1				10.400
339	HIV-1	0.123	83.49				HIV1TANDEM				5.300
	HIV-1	0.370	100.6				HIV-1	0.123		51.83	
	HIV-1	1.100	101.42				HIV-1	0.370		68.49	
	HIV-1	3.300	104.71		35		HIV-1	1.100		70.71	
	HIV-1	10.000	91.38				HIV-1	3.300		63.96	
	HIV-1	30.000	72.86				HIV-1	10.000		51.8	
	HIV1TANDEM			0.120			HIV-1	30.000		43.93	
340	HIV-1	0.123	80.58			354	HIV-1	0.123	<	10	
	HIV-1	0.370	90.49				HIV-1	0.370		10.37	
	HIV-1	1.100	90.16		40		HIV-1	1.100		26.79	
	HIV-1	3.300	91.57				HIV-1	3.300		46.1	
	HIV-1	10.000	89.49				HIV-1	10.000		54.97	
	HIV-1	30.000	71.99	0.060			HIV-1	30.000		54.5	CCE 000
242	HIV1TANDEM	0.100	01.06	0.060		255	HIV-1				665.000
342	HIV-1	0.123	81.06			355	HIV-1	0.122		10	700.000
	HIV-1	0.370	93.18		45		HIV-1	0.123 0.370	<	10	
	HIV-1	1.100 3.300	96.94 85.55				HIV-1 HIV-1	1.100	<	10	
	HIV-1 HIV-1	10.000	73.55				HIV-1	3.300	<	20.72	
	HIV-1	30.000	73.95				HIV-1	10.000		46.66	
	HIV1TANDEM	30.000	13.33	0.309			HIV-1	30.000		67.82	
343	HIV-1	0.123	57.5	0.509		356	HIV1TANDEM	50.000		07.02	1.100
545	HIV-1	0.370	76.83		50	330	HIV-1	0.123		54.96	1.100
	HIV-1	1.100	81.02		30		HIV-1	0.370		71.75	
	HIV-1	3.300	86.43				HIV-1	1.100		90.19	
	HIV-1	10.000	60.56				HIV-1	3.300		92.28	
	HIV-1	30.000	46	2.900			HIV-1	10.000		100.22	
344	HIV-1	0.123	47.37				HIV-1	30.000		95.16	
•	HIV-1	0.370	72.84		55	357	HIV1TANDEM				48.500
	HIV-1	1.100	81.17		33	359	HIV1TANDEM				16.400
	HTV-1	3.300	83.08			363	HIV1TANDEM				0.083
	HIV-1	10.000	68.47			365	HIV1TANDEM				0.023
	HIV-1	30.000	46.24	5.900		368	HIV1TANDEM				0.232
345	HIV1TANDEM			0.032		370	HIV-1	0.123		92.81	
	HIV-1	0.123	69.19		<b>C</b>		HIV-1	0.370		87.87	
	HIV-1	0.370	94.37		60		HIV-1	1.100		102.89	
	HIV-1	1.100	101.67				HIV-1	3.300		109.33	
	HIV-1	3.300	99.08				HIV-1	10.000		113.79	
	HIV-1	10.000	97.43				HIV-1	30.000		98.14	0.590
	HIV-1	30.000	84.56				HIV1TANDEM				0.050
347	HIV1TANDEM			13.600			HIV-2				0.050
	HIV-1	0.123	20.99		65	371	HIV-1	0.123		39.84	
	HIV-1	0.370	50.82				HIV-1	0.370		72.94	

TABLE II-continued

		TABL	E II-contin	ued						TABLE 3-continued	
EXAMI	PLE SER EN	ZYMF	DOSE	PROTE % INH		FITC KI (NM)		U-No.	MS data	Name	Origin
NOND	HIV	7-1 7-1 7-1	1.100 3.300 10.000 30.000	91.0 104.1 102.1 107.1	61 12 7	15.400	. 5	309	565.2605 (EI)	N-[3(R or S)-[1-(5,6-Dihydro-4- hydroxy-2-oxo-6(R or S)-[2- phenylethyl]-6-propyl-2H-pyran-3- yl)-2,2-dimethylpropyl]phenyl]-1- methyl-1H-imidazole-4-	Single stereoisomer; Derived from Isomer 3 of Preparation 143
372	HIV HIV HIV HIV	/-1 /-1 /-1 /-1	0.123 0.370 1.100 3.300 10.000 30.000	43.5 86.6 101.5 99.5 97.8 106.1	68 52 56 31	4.000	10	310	565.2626 (EI)	sulfonamide N-[3(R or S)-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6(R or S)-[2-phenylethyl]-6-propyl-2H-pyran-3-yl)-2,2-dimethylpropyl]phenyl]-1-methyl-1H-imidazole-4-	Single stereoisomer; Derived from Isomer 4 of Preparation 148
373	HIV HIV HIV HIV HIV	7-1 7-1 7-1 7-1	0.123 0.370 1.100 3.300 10.000 30.000	90.3 90.3 103.8 88.3 85.3 89.5	35 33 72 75		15	311	571.2113 (EI)	sulfonamide 5-Cyano-N-[3-(1-[5,6-dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]cyclopropyl-methyl)phenyl]-2-pyridine-sulfonamide	Diastereomeric mixture
374	HIV	71TANDEM 71TANDEM	0.123 0.370 1.100 3.300 10.000 30.000	78.9 82.1 84.9 87.7 95.2 80.1	97 14 98 70 25	0.200	20	312	577.2630 (EI)	5-Amino-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]-2,2-dimethylpropyl)-phenyl]-2-pyridinesulfonamide	Single stereoisomer; Prepared from amine of Preparation 138 (derived from Isomer 1 of Preparation
		ī	ABLE 3				25	313	577.2585 (EI)	5-Amino-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-	143) Single stereoisomer; Prepared from
U-No.	MS data	Name			Origi	n	30			pyran-3-yl]-2,2-dimethylpropyl)- phenyl]-2-pyridinesulfonamide	amine of Preparation 137 (derived from
	587.2453 (EI)	dihydro-4-l S)-(2-phen	I-[3(R or S)-(1 hydroxy-2-oxo ylethyl)-6-prop J-2,2-dimethylp	-6(R or yl-2H-	Deriv	e pisomer; red from er 1 of		314	550.2380	5-Amino-N-[3(R or S)-(1-[5,6-	Isomer 2 of Preparation 143) Single
	587.2458 (EI)	5-Cyano-N dihydro-4-l S)-(2-phen pyran-3-yl]	oyridinesulfons -[3(R or S)-(1- hydroxy-2-oxo ylethyl)-6-prop  -2,2-dimethylp	-[5,6- -6(R or oyl-2H- oropyl)-	Single stereo Deriv Isome	oisomer; ed from er 2 of	35	315	(FAB) 550.2365	dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl)phenyl]-2-pyridinesulfonamide 5-Amino-N-[3(R or S)-(1-[5,6-	stereoisomer; Derived from Isomer 1 of Preparation 147 Single
	587.2444 (EI)	5-Cyano-N dihydro-4-l S)-(2-phen	oyridinesulfons -[3(R or S)-(1- hydroxy-2-oxo ylethyl)-6-prop  -2,2-dimethylp	-[5,6- -6(R or -yl-2H-	Single stereo Deriv	ration 143 e visomer, ed from er 3 of	40	316	(FAB) 596.2583	dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl)phenyl]-2-pyridinesulfonamide 5-Amino-N-[3(R or S)-(1-[6(R or	stereoisomer; Derived from Isomer 2 of Preparation 147 Single
	587.2446 (EI)	5-Cyano-N dihydro-4-l S)-(2-pheny	oyridinesulfona -[3(R or S)-(1- hydroxy-2-oxo- ylethyl)-6-prop  -2,2-dimethylp	-{5,6- -6(R or -yl-2H-	Single stereo Deriv	ration 143 e isomer, ed from er 4 of	45		(FAB)	S)-(2-[4-fluorophenyl)ethyl)-5,6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethyl propyl)phenyl]-2-pyridinesulfonamide	stereoisomer; Derived from Isomer 1 of Preparation 150
	525.2311 (EI)	phenyl)-2-r 5-Cyano-N diisobutyl-	oyridinesulfons -[3-(1-[5,6-dih 4-hydroxy-2-ox  propyl)phenyl	mide ydro-6,6- ro-2H-		ration 143 nic	•	317	596.2583 (FAB)	5-Amino-N-[3(R or S)-(1-[6(R or S)-(2-[4-fluorophenyl]ethyl)-5,6-dihydro-4-hydroxy-2-oxo-6-propyl-2H-pyran-3-yl]-2,2-dimethyl-propyl)phenyl]-2-pyridine-	Single stereoisomer; Derived from Isomer 2 of Preparation 150
(	532.2856 (FAB)	diisobutyl-4 pyran-3-yl] phenyl]-1-r sulfonamid	6-Dihydro-6,6- 4-hydroxy-2-oz  -2,2-dimethylp nethyl-1H-imic e	ro-2H- ropyl)-	Racen		50	318	503.2445 (EI)	sulfonamide N-[3(R or S)-(1-[5,6-Dihydro-6,6- dipropyl-4-hydroxy-2-oxo-2H- pyran-3-yl]-2,2-dimethylpropyl)- phenyl]-1-methyl-1H-imidazole-4-	Single enantiomer, Prepared from amine derived
	554.2688 (FAB)	6-diisobuty pyran-3-yl]	-[3-(1-[5,6-dih l-4-hydroxy-2- -2,2-dimethylp pyridinesulfona	oxo-2H- ropyl)-	Racen mixtu		55	319	502 2454	sulfonamide  N-[3(R or S)-(1-[5,6-Dihydro-6,6-	from Isomer 1 of Preparation 144
	565.2607 (EI)	N-[3(R or ship) hydroxy-2- phenylethyl yl)-2,2-dim	S)-[1-(5,6-Dihy oxo-6(R or S)- l]-6-propyl-2H ethylpropyl]ph -imidazole-4-	ydro-4- [2- -pyran-3-	Derive Isome	isomer; ed from	60	313	503.2454 (EI)	N-13(k of 5)-(1-13,6-Dinydro-6,6-dipropyl-4-hydroxy-2-0xo-2H-pyran-3-yl]-2,2-dimethylpropyl)-phenyl]-1-methyl-1H-imidazole-4-sulfonamide	Single enantiomer; Prepared from amine derived from Isomer 2 of Preparation 144
	565.2629 (EI)	N-[3(R or 5 hydroxy-2- phenylethyl yl)-2,2-dim	S)-[1-(5,6-Dihy oxo-6(R or S)- l}-6-propyl-2H ethylpropyl]ph imidazole-4-	[2- -pyran-3- enyl]-1-	Derive Isome	isomer; ed from	65	320	515.2453 (EI)	5-Amino-N-[3(R or S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethylpropyl)phenyl]-2-pyridinesulfonamide	Single enantiomer, Prepared from amine derived from Isomer 1 of Preparation

TABLE 3-continued

U-No.	MS data	Name	Origin	
321	515.2463 (EI)	5-Amino-N-[3(R or S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethyl-propyl)phenyl]-2-pyridine-sulfonamide	144 Single enantiomer; Prepared from amine derived from Isomer 2 of Preparation 144	10
322	525.2287 (EI)	5-Cyano-N-[3(R or S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethyl-propyl)phenyl]-2-pyridine-sulfonamide	Single enantiomer; Prepared from amine derived from Isomer 2 of Preparation 144	15
323	525.2288 (EI)	5-Cyano-N-[3(R or S)-(1-[5,6-dihydro-6,6-dipropyl-4-hydroxy-2-oxo-2H-pyran-3-yl]-2,2-dimethyl-propyl)phenyl]-2-pyridinesulfonamide	Single enantiomer; Prepared from amine derived from Isomer 1 of Preparation 144	20
324	600.2537 (FAB)	N-[3(R or S)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide	Single enantiomer, Prepared from amine derived from Isomer 1 of Preparation 145	25
325	600.2537 (FAB)	N-[3(R or S)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-1-methyl-1H-imidazole-4-sulfonamide	Single enantiomer, Prepared from amine derived from Isomer 2 of Preparation 145	30
326	622.2378 (FAB)	N-[3(R or S)-(1-[6,6-Bis(2-phenyl- ethyl)-5,6-dihydro-4-hydroxy-2- oxo-2H-pyran-3-yl]propyl)phenyl]- 5-cyano-2-pyridinesulfonamide	Single enantiomer; Prepared from amine derived Isomer 1 of Preparation 145	35
327	622.2367 (FAB)	N-[3(R or S)-(1-[6,6-Bis(2-phenyl- ethyl)-5,6-dihydro-4-hydroxy-2- oxo-2H-pyran-3-yl]propyl)phenyl]- 5-cyano-2-pyridinesulfonamide	Single enantiomer; Prepared from amine derived from Isomer 2 of Preparation 145	40

We claim:

### 1. The compound of the formula VI

$$\begin{array}{c} OH & VI \\ R_2 & \\ R_3 & O \end{array}$$

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x 60

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wherein  $R_2$  is a)  $H_3C$ — $CH_2$ —, or

b) phenyl- $(CH_2)_2$ —;

wherein R<sub>3</sub> is the moiety of formula X

wherein R6 is a)  $H_3C-(CH_2)_2$ —, or b) phenyl-(CH<sub>2</sub>)<sub>2</sub>--;

wherein R<sub>7</sub> is H<sub>3</sub>C—CH<sub>2</sub>—;

wherein R<sub>9</sub> is ---NHSO<sub>2</sub>---het;

wherein het is 2-pyridinyl substituted at the 5-position by zero (0) or one (1) R<sub>10</sub>;

wherein R<sub>10</sub> is

a) —CN,

b) —CF<sub>3</sub>,

c) —NH2, or

d) -CONH2;

or a pharmaceutically acceptable salt thereof.

2. The compound of claim 1 selected from the group consisting of:

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H<sub>3</sub>C

H₃Ċ

$$H_3C$$
 $OH$ 
 $CH_3$ 
 $C=N$ 
 $C=N$ 

H<sub>3</sub>C OH CH<sub>3</sub> 
$$C \equiv N$$

- 3. The compound of claim 1 selected from the group consisting of:
  - 5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide;
  - 5-Trifluoromethyl-N-[3-(R)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide; or (3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl] phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
  - 5-Trifluoromethyl-N-[3-(S)-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide; or (3S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl] phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- 5-Trifluoromethyl-N-[3(R or S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenethyl)-6(R or S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridine sulfonamide:
- 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -propyl]-phenyl]-2-pyridinesulfonamide; or (3R,6R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl)propyl]-phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- 5-Trifluoromethyl-N-[3(R)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide; or (3R,6S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6.(2-phenylethyl)-6-propyl-2H-pyran-3-yl)propyl]-phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- 5-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(R)-(2-phenethyl)-6(R)-n-propyl-2H-pyran-3-yl] -propyl]-phenyl]-2-pyridinesulfonamide; or (3S,6R)-N-[3-[1-[5,6,-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- 5-Trifluoromethyl-N-[3(S)-[1-[5,6-dihydro-4-hydroxy-2-oxo-6(S)-(2-phenethyl)-6(S)-n-propyl-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide; or (3S,6S)-N-[3-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- N-[3-[1-(S)-[5,6,-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- (3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-sulfonamide;

- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R or S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- N-[3-{1(R)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(R)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5- 15 cyanopyridine-2-sulfonamide;
- N-[3-{1(S)-(4-Hydroxy-5,6-dihydro-2-oxo-6(S)-phenethyl-6-propyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- 5-Amino-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl] propyl)phenyl]-2-pyridinesulfonamide;
- 5-Amino-N-[3(R or S)-(1-[5,6-dihydro-4-hydroxy-2-oxo-6(R or S)-(2-phenylethyl)-6-propyl-2H-pyran-3-yl] propyl)phenyl]-2-pyridinesulfonamide;
- 5-Trifluoromethyl-N-[3-[1-[4-hydroxy-2-oxo-6,6-di-n-propyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide;
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-30 phenylethyl)-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide;
- N-[3(R or S)-(1-[6,6-Bis(2-phenylethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide;
- N-[3(R or S)-(1-[6,6-Bis(2-phenyl-ethyl)-5,6-dihydro-4-hydroxy-2-oxo-2H-pyran-3-yl]propyl)phenyl]-5-cyano-2-pyridinesulfonamide;
- N-[3-{1(R or S)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-aminopyridine-2-sulfonamide;
- N-[3-{1(S or R)-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)-propyl}phenyl]-5-aminopyridine-2-sulfonamide;

- 5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-phenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide;
- 5-Trifluoromethyl-N-[3-(R or S)-[1-[4-hydroxy-2-oxo-6, 6-di-phenethyl-5,6-dihydro-2H-pyran-3-yl]-propyl]-phenyl]-2-pyridinesulfonamide;
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6-phenethyl-6propyl-2H-pyran-3-yl)propyl}phenyl]-5cyanopyridine-2-sulfonamide;
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl}phenyl]-5-cyanopyridine-2-sulfonamide; and
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-dipropyl-2H-pyran-3-yl)propyl}phenyl]-5-carbamoylpyridine-2-sulfonamide.
- 4. The compound of claim 1 selected from the group consisting of:
  - (3R,6R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
  - (3R,6S)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- (3S,6R)-N-[3-[1-[5,6,-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- (3S,6S)-N-[3-[1-[5,6-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide;
- N-[3-[1-(S)-[5,6,-Dihydro-4-hydroxy-2-oxo-6-(2-phenylethyl)-6-propyl-2H-pyran-3-yl]propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide; and
- (3R)-N-[3-[1-(5,6-Dihydro-4-hydroxy-2-oxo-6-propyl-6-phenethyl-2H-pyran-3-yl)propyl]phenyl]-5-(trifluoromethyl)-2-pyridinesulfonamide.
- 5. A compound selected from the group consisting of:
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)methyl}phenyl]-5-cyanopyridine-2-sulfonamide, and
- N-[3-{1-(4-Hydroxy-5,6-dihydro-2-oxo-6,6-bis(2-phenylethyl)-2H-pyran-3-yl)methyl}phenyl]-5-aminopyridine-2-sulfonamide.

\* \* \* \* \*

EXHIBIT C COPY OF CERTIFICATE OF CORRECTION FOR U.S. PATENT 5,852,195

# UNITED STATES PATENT AND TRADEMARK OFFICE CERTIFICATE OF CORRECTION

PATENT NO.

: 5,852,195

: December 22, 1998

DATED INVENTOR(S)

INVENTOR(S) : Romines, et al.

hereby corrected as shown below:

It is certified that error appears in the above-identified patent and that said Letters Patent is

Column 389,

Lines 60-65:

wherein R<sub>6</sub> is

a) $H_3C$ —( $CH_2$ )<sub>2</sub>—, or

Signed and Sealed this

Page 1 of 1

Ninth Day of October, 2001

Anest:

Nicholas P. Ebdici

Attesting Officer

NICHOLAS P. GODICI
Acting Director of the United States Patent and Trademark Office

EXHIBIT D COPY OF MAINTENANCE FEE RECORDS









## **Maintenance Fee Statement**

07/08/2005 06:31 F

Patent Number: 5852195

Customer Number: 000000

MARTHA A. GAMMILL INTELLECTUAL PROPERTY LEGAL SERVICES PHARMACIA AND UPJOHN COMPANY KALAMAZOO MI 49001

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PATENT NUMBER	FEE AMT	SUR- CHARGE	U.S. APPLICATION NUMBER	PATENT ISSUE DATE	APPL. FILING DATE	PAYMENT YEAR	SMALL ENTITY?	STAT
5,852,195	\$880.00	\$0.00	08/809,224	12/22/98	11/04/96	04	NO	PAID

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Patent Mainte	nance Fees	07/08/2005 06:33 PM EDT			
Patent Number:	5852195	<b>Application Number:</b>	08809224		
Issue Date:	12/22/1998	Filing Date:	11/04/1996		
Window Opens:	12/22/2005	Surcharge Date:	06/23/2006		
Window Closes:	12/22/2006	Payment Year:			
Entity Status:	LARGE				
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EXHIBIT E
INFORMATION TO ENABLE DETERMINATION
OF REGULATORY REVIEW PERIOD

#### Relevant Dates and Information to Enable Determination of the Regulatory Review Period

The '195 patent claims a human drug product.

The Investigational New Drug (IND) application for the APTIVUS® product, IND No. 51979, was submitted on November 13, 1996, received by the FDA on November 14, 1996, and became effective 30 days after submission on December 13, 1996, pursuant to 21 C.F.R. §312.

The New Drug Application (NDA) for the APTIVUS® product, NDA No. 21-814, was submitted on December 21, 2004 and was approved by the FDA on June 22, 2005.

#### **EXHIBIT F**

#### **DESCRIPTION OF SIGNIFICANT ACTIVITIES**

This exhibit includes the following four chronolgies:

- Agency Contact Reports Any phone calls, meetings or emails between BI and FDA
- Correspondence from FDA Faxes or letters (hard copy) from FDA to BI
- IND Log Listing of all submissions to the tipranavir IND 51,979
- NDA Log Listing of all amendments to the tipranavir NDA 21-814 (NDA 21-822 (solution) cross references 21-814 for all clinical and non-clinical data)

Description of Reference			
Serial	Number	Jo	Reference
Date of	Reference		
Description of FDA Correspondence			
Date of FDA	Correspondence		

11/21/96	FDA Letter Assigning IND Number	11/13/96	000	Original IND Application
12/17/96	FDA Fax providing clinical, pharmacology and	11/13/96	000	Original IND Application
	Protocol M/3342/001	12/13/96	N/A	PN&U and FDA Teleconference
		12/13/96	001	Protocol Amendment – Change in Protocol M/3342/0001
12/19/96	FDA Fax requesting information on drug substance drug product and biopharmaceutics	11/13/96	000	Original IND Application
	on New IND and Protocol M/3342/001	12/13/96	N/A	PN&U and FDA Teleconference
		12/13/96	001	Protocol Amendment - Change in Protocol M/3342/0001
1/27/97	FDA Letter providing clinical, pharmacology and microbiology comments on the IND and	12/13/96	N/A	PN&U and FDA Teleconference
	Protocol M/3342/001	12/17/96	N/A	FDA Fax providing clinical, pharmacology and microbiology comments on New IND
	Request for information regarding drug substance, drug product, and biopharmaceutics			and Protocol M/3342/001
		12/19/96	N/A	FDA Fax requesting information on drug substance, drug product and biopharmaceutics on New IND and Protocol M/3342/001
76/11/7	Letter informing PNU about the AIDS Clinical Trials Information Service (ACTIS) and requesting that PNU submits information about Protocol M/3342/0004 for the inclusion for the	1/2/97	003	Protocol Amendment – Change in Protocol M/3342/001 Amendment 2
	ACTIS database			

Description of Reference	
Serial Number	of Reference
Date of Reference	
Description of FDA Correspondence	
Date of FDA Correspondence	

2/27/98	FDA Fax providing biopharmaceutics comments on draft protocol M/3342/0009	2/4/98	018	Protocol Amendment – New Protocol Draft M/3342/0009
9/24/98	FDA Fax providing Biopharmaceutics comments on Protocol M/3342/0012	8/25/98	028	Protocol Amendment – New Protocol M/3342/0018
9/27/98	FDA Fax providing biopharmaceutics comments on Protocol M/3342/0009	2/4/98	018	Protocol Amendment – New Protocol Draft M/3342/0009
		2/27/98	810	FDA Fax providing biopharmaceutics comments on draft protocol M/3342/0009
3/24/99	FDA Request for information – Clinical Comments on SN 040 and SN 041	1/29/99	040	Annual Report – Reporting Period 10/1/97 – 9/30-98
·		2/5/99	041	Protocol Amendment – New Protocol M3342/0006
4/21/99	Letter informing PNU about the AIDS Clinical Trials Information Service (ACTIS) and requesting that PNU submits information about Protocol M/3342/0006 for the inclusion for the	2/5/99	<u>4</u>	Protocol Amendment – New Protocol M/3342/0006
	ACTIS database	3/29/99	043	Protocol Amendment –  New Protocol M/3342/0013  Change in Protocol M/3342/0006  Amendment 1
5/19/99	FDA Fax providing the FDA Meeting minutes from the May 11, 1999 Meeting discussing the Rat Carcinogenicity Study Dose Selection	5/11/99	N/A	FDA/PNU Meeting discussing the Rat Carcinogenicity Study Dose Selection

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Description of Reference				
Serial	Number	of	Reference	
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Description of FDA Correspondence				
Date of FDA	Correspondence			

5/21/99	Letter informing PNU about the AIDS Clinical Trials Information Service (ACTIS ) and requesting that PNU submits information about Protocol M/3342/0016 for the inclusion for the ACTIS database	5/10/99		Protocol Amendment –  • New Protocol M/3342/0016
6/2/99	FDA Fax containing clinical comments on SN 049	5/10/99	049	<ul> <li>Protocol Amendment –</li> <li>New Protocol M/3342/0016</li> <li>New Investigator for Protocol M/3342/0006</li> <li>New Investigator for Protocol M/3342/0013</li> <li>New Investigator for Protocol 69INF0013-019</li> <li>Change in Investigator for Protocol M/3342/0015</li> </ul>
9/15/99	Letter encouraging all sponsors with antiretroviral drugs in development to conduct studies in treatment-exposed HIV-infected individuals and to promote pharmaceutical collaboration to meet this objective	N/A	N/A	N/A
5/15/00	Letter from FDA acknowledging transfer of IND from Pharmacia & Upjohn to Boehringer Ingelheim Pharmaceuticals effective April 21, 2000	4/19/00	890	General Correspondence Transfer of Sponsorship from Pharmacia & Upjohn Company to Boehringer Ingelheim Pharmaceuticals, Inc.
			690	General Correspondence – Boehringer Ingelhe im's acknowledgment of transfer of IND Sponsorship

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Description of FDA Correspondence			
Date of FDA	Correspondence		

									S.				<u> </u>				
Original Protocol Amendment 1	Amendment 2 and 3	Amendment 4 and 5	Amendment A, 6 and 7	Amendment 8	Amendment 9	Amendment 10	Amendment 11	Amendment 12	Protocol Amendment - New Protocol 1182.5		Protocol Amendment - New Protocol and	New Investigator for 1182.6	Protocol Amendment - New Protocol and	New Investigator for 1182.6	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	FLOA TAX PROVIDING CHINICAL AND	1182.6
010 012	015	018	022	025	027	039	029	071	072		9/0		9/0		720	0/0	
79/10/77 8/22/97	9/30/97	2/4/98	4/1/98	86/2/9	86/08//	12/9/98	11/23/99	8/3/00	9/13/00		1/19/01		1/19/01		3/1/01	3/1/01	
Fax including FDA comments on Amendment 12 of Trial No. 1182.1									Fax containing FDA comments on Protocol	1182.5	FDA fax providing clinical and	pharmacokinetics comments on Protocol 1182.6	FDA fax providing microbiological comment on	Protocol 1182.6			
9/13/00									11/9/00		3/1/01		3/5/01				-

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3/19/01	FDA Fax with clinical comments and requests	3/19/01	8/0	Protocol Amendment
	for information regarding BI Trials			<ul> <li>New Protocol for 1182.17</li> </ul>
				<ul> <li>Change in Protocol 1182.4</li> </ul>
				Amendment 2
				<ul> <li>Change in Protocol 1182.6</li> </ul>
				Amendment 1
6/29/01	Official FDA Meeting Minutes from the End of Phase I Meeting on April 5, 2001	4/5/01	220	End of Phase I Background Document
11/15/01	FDA Fax containing comments on SN 122	11/7/01	122	General Correspondence - Request for
				Advice on Duration of Combination Toxicology Studies
11/16/01	FDA Fax with clinical comments on BI Trial	11/16/01	121	Protocol Amendment
	1182.37 and 1182.41			<ul> <li>New Protocols 1182.37 and 1182.41</li> </ul>
				<ul> <li>Changes in Protocol 1182.37</li> </ul>
				Amendment 1
11/30/01	Official FDA Meeting Minutes from Clinical	10/2/01	N/A	Meeting to discuss clinical development plans
	Development plans and drug interaction program for tipranavir on October 5, 2001			and drug interaction program
		11/7/01	123	BIPI's Meeting Minutes from 10/5/01 meeting
12/6/01		11/16/01	121	Protocol Amendment
	on BI Trial 1182.41		-	<ul> <li>New Protocols 1182.37 and 1182.41</li> </ul>
2/4/02	Fax from FDA requesting additional	12/27/01	133	Response to FDA Comments in regards to
	information in reference to SN 133			combination toxicology studies

Description of Reference		
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Description of FDA Correspondence		
Date of FDA Correspondence		

70/01/7	Fax from FDA Requesting Information from BI	1/28/02	132	Information Amendment – Clinical
	in order to address the clinical development program. FDA Requesting Chemistry,			Pharmacokinetics from 1182.6
	Pharmacology/Toxicology, Pharmacokinetics and Clinical information.	2/4/02	N/A	Fax from FDA Requesting additional Information on SN 133
		12/27/01	133	Response to FDA Comments in regards to combination toxicology studies
3/22/02	Letter encouraging all sponsors with	N/A	N/A	N/A
	antiretroviral drugs in development to conduct studies in treatment-exposed HIV-infected			
	individuals and to promote pharmaceutical			
	collaboration to meet this objective ACTIS	٠		
	database.			
4/26/02	Letter encouraging all sponsors with	N/A	N/A	N/A
	antiretroviral drugs in development to conduct			
	studies in treatment-exposed HIV-infected			
	individuals and to promote pharmaceutical			
	collaboration to meet this objective			
5/17/02	Letter encouraging all sponsors with	N/A	N/A	N/A
	antiretroviral drugs in development to conduct			
	studies in treatment-exposed HIV-infected			
-	individuals and to promote pharmaceutical			
	collaboration to meet this objective			

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Description of FDA Correspondence			7 W M M M M M M M M M M M M M M M M M M
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7/24/02	FDA Fax with Pharmacology comments on SN 162; FDA agrees with proposed change of dose level from 26.7 mg/kg/day for the ritonavir only arm of the rat carcinogenicity study protocol	7/19/02	162	Information Amendment- Pharmacology/Toxicology Reports     Request for FDA Feedback/Teleconference – BI request concurrence with lowering dose level of ritonavir    Page 12
8/29/02	Email from FDA with copy of FDA letter dated 8/29/02 acknowledging receipt of special clinical protocol assessment. FDA believes	7/18/02	191	Request for Special Protocol Assessment- Clinical BI requests special protocol assessment of 1182.52
	submission is not adequate for complete special protocol assessment because data is missing. FDA provides information that is required and refers to guidance for additional information.	6/7,6/18, 6/19 and 6/25/02	ACR	Feedback on Special Protocol Assessment
8/29/02	FDA letter with acknowledging receipt of special clinical protocol assessment. FDA believes submission is not adequate for complete special protocol assessment because	7/18/02	161	Request for Special Protocol Assessment- Clinical BI requests special protocol assessment of 1182.52
	data is missing. FDA provides information that is required and refers to guidance for additional information.	8/29/02	161	Email from FDA providing FDA Letter
9/20/02	FDA Fax providing clinical pharmacology comments on SN 137. FDA believes BI has not submitted sufficient information for the Division to conclude that further investigation	2/14/02	137	Information Amendment: Clinical/Request for Comment-Discontinuation of Tipranavir Study 1182.42
	of the drug interaction potential between ddl and tipranavir/ritonavir (TPV/RTV) is unnecessary. Reasons are provided.	12/20/01	132	Information Amendment - Clinical Pharmacokinetics from 1182.6

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		g.		
Request for Special Protocol Assessment- Clinical BI requests special protocol assessment of 1182.52	Request for Special Protocol Assessment- Clinical BI requests special protocol assessment of 1182.52	FDA letter with acknowledging receipt of special clinical protocol assessment. FDA believes submission is not adequate for complete special protocol assessment because data is missing. FDA provides information that is required and refers to guidance for additional information.	Request for Special Protocol Assessment- Clinical BI requests special protocol assessment of 1182.12	Fax from FDA with Comments on SN 161. FDA provides answers to the questions that were included in the Request for Special Protocol Assessment. Additional comments are also provided.
161	191	X/X	161	N/A
7/18/02	7/18/02	8/29/02	7/18/02	9/20/02
		<b>b</b> 0 0		6
Fax from FDA with Comments on SN 161. FDA provides answers to the questions that were included in the Request for Special Protocol Assessment. Additional comments are also provided.	Letter encouraging all sponsors with antiretroviral drugs in development to conduct studies in treatment-exposed HIV-infected individuals and to promote pharmaceutical collaboration to meet this objective	FDA Fax with Overview of Pharmacology/Toxicology comments regarding the clinical development of tipranavir  Human exposure  Justification of Formulation (including info on vehicle	-	
9/20/02	10/3/02	10/30/02		

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		3/5/02- 5/17/02	139 150	Response to FDA Request - Providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials
12/10/02	Fax from FDA with Clinical and Clinical Pharmacology comments regarding SN 186. FDA Requests that responses are provided prior to the End of Phase II Meeting	11/15/02	186	General Correspondence - Background Document for End of Phase II Meeting
12/12/02	FDA Fax with Pharmacology/Toxicology comments in preparation for the December 12, 2002 teleconference	12/4/02	N/A	Teleconference between BI and FDA
12/31/02	FDA Fax providing Clinical and Statistical comments regarding Protocol 1182.12	11/15/02	186 N/A	General Correspondence - Background Document for End of Phase II Meeting End of Phase II Meeting

Description of Reference			
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Description of FDA Correspondence			
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1/3/03	FDA Fax with Microbiology comments regarding SN 186	11/15/02	186	General Correspondence - Background Document for End of Phase II Meeting
		12/17/02	N/A	End of Phase II Meeting
1/14/03	FDA Official Meeting Minutes from the End of Phase II Meeting	12/17/02	N/A	End of Phase II Meeting
		3/14/03	N/A	FDA End of Phase II Meeting Minutes
		3/24/03	217	BIPI End of Phase II Meeting Minutes
1/22/03	FDA Letter requesting additional information	11/15/02	186	General Correspondence - Background
	on pediatric studies. BI must submit the			Document for End of Phase II Meeting
	enterestion mitting 100 deep	2714/03	A1/A	FDA F. 4 - CDL HAK C AK'
	submissions within 180 days.	3/14/03	A/Z	FDA End of Phase II Meeting Minutes
		3/24/03	217	BIPI End of Phase II Meeting Minutes
4/14/03	FDA's response to New Protocol 1182.51	1/30/03	204	Protocol Amendment - New Protocol 1182.51
5/7/03	FDA Fax providing Response to	3/24/03	216	Request for Special Protocol Assessment
	Carcinogenicity Special Protocol Assessment			Carcinogenicity Study Protocol. BI asking
	Report - Final CAC Report. FDA recommends			FDA concurrence with the mouse
	BI to evaluate the dosage groups by pair wise			carcinogenicity study protocol.
	companion with the inclusive control.	20/2/9	ACRS	Special Protocol Assessment for mouse
		6/18/02		carcinogenicity protocol.
	٠.	6/19/02 6/25/02		
5/30/03	FDA Approval to export tipranavir to Brazil for	5/12/03	N/A	Request to Export Drug to Brazil for 1182.48
		5/8/03	ACR	DAVDP's review of export
				waiver
		4/8/03 -	ACR	Export waiver status

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		11/03		
5/30/03	FDA Approval to export tipranavir to Argentina for 1182.48	3/12/04	N/A	Request to Export Drug to Argentina for 1182.48
		4/8/03- 11/03	ACR	Export waiver status
5/30/03	FDA Approval to export tipranavir to Mexico for 1182.48	3/14/04	N/A	Request to Export Drug to Mexico for 1182.48
9/4/03	Response to BI's request for a teleconference to discuss proposed pediatric written agreement.  FDA classified the meeting as Type B and it is	7/23/03	242	Proposed Changes in Written Request for Pediatric Studies
	scheduled for September 18, 2003	8/26/03 8/15/03	ACR ACR	Proposed date of telecom Request for telecon
		1/22/03 & 1/28/03	N/A	Proposed Written Agreement for Pediatric Studies
9/15/03	for	7/23/03	N/A	Request to Export Drug to Brazil for 1182.17
9/16/03	FDA Fax providing statistical comments on Protocol 1182.12 (RESIST 1)	3/19/03	SN 214	Protocol Amendment – Changes in Protocol 1182.12 Amendment 1 and 2
		2/4/03	SN 205	Original Protocol

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Description of FDA Correspondence			
Date of FDA	Correspondence		

9/17/03	FDA Approval to export tipranavir to Argentina for 1182.17	7/23/03 6/6/03	N/A ACR	Request to Export drug to Argentina for 1182.17
		9/6/03-9/8/03 ACR	ACR	Export waiver – 1182.17 for Argentina
		8/23/03- 8/29/03	ACR	Request for status of export waiver
		8/6/03-8/8/03 ACR	ACR	Request for status of export waiver
		8/12/03- 8/15/03	ACR	Request for status of export waiver
		7/31/03- 8/15/03	ACR	Request for status of export waiver
9/25/03	FDA Fax containing Comments regarding responses to proposed written agreement for pediatric studies	7/23/03	242	Proposed Changes in Written Request for Pediatric Studies Proposed Written Agreement for Pediatric
				Studies Request for Teleconference to discuss submission
10/31/03	FDA Official Meeting Minutes from the teleconference discussing the Proposed Changes in Written Request for Pediatric Studies	10/2/03	N/A	BI and FDA Teleconference discussing the Proposed Changes in Written Request for Pediatric Studies
		7/23/03	186	Proposed Changes in Written Request for Pediatric Studies

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11/13/03	FDA Letter with Responses to Request for Special Protocol Assessment – Clinical, Naïve Trial 1182, 33	9/26/03	254	Request for Special Protocol Assessment – Clinical, Naïve Trial 1182.33
		10/24/03	ACR	Naïve Trial 1182.33
		7/18/03	ACR	Naïve Trial 1182.33
		7/15/03	ACR	Naïve Trial 1182.33
		12/17/02	N/A	End of Phase II Meeting
11/13/03	FDA Acknowledgment of Receipt of Request	9/26/03	254	Request for Special Protocol Assessment -
	for Special Protocol Assessment – Clinical,			Clinical, Naïve Trial 1182.33
	1/a1/C 111a1 1102.33			
11/14/03	FDA Approval to export tipranavir to Brazil for 1182.14	8/20/03	N/A	Request to Export Drug to Brazil for 1182.14
11/14/03	FDA Approval to export tipranavir to Argentina 8 for 1182.14	8/20/03	N/A	Request to Export Drug to Argentina for 1182.14
11/14/03	FDA Approval to export tipranavir to Mexico for 1182.14	8/20/03	N/A	Request to Export Drug to Mexico for 1182.14
	The state of the s			
11/14/03	FDA Approval to export tipranavir to Russia for 1182.14	9/8/03	N/A	Request to Export Drug to Russia for 1182.14

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Description of FDA Correspondence			
Date of FDA	Correspondence		

12/12/03	FDA Approval to export tipranavir to Russia,	60/8/63	N/A	Request to Export Drug to Russia for
	Diazii, Augeliulia aliu Mealeo 101 1102.14	8/20/03	N/A	1102.14
		60/06/0	****	Request to Export Drug to Brazil for 1182.14
		\$/70/03	N/A	Domest to Evnet Dens to Arganting for
				request to export orug to Augentina for 1182.14
		8/20/03	N/A	
				Request to Export Drug to Mexico for 1182.14
2/20/04	Letter from FDA confirming with BI that a	1/30/04	297	General Correspondence - Request for Pre-
	Type B Meeting is scheduled for May 10, 2004.			NDA Meeting
3/3/04	FDA Official Meeting Minutes from the	1/16/04	N/A	Meeting discussing BI's safety reporting
	Meeting discussing BI's safety reporting policies and procedures			procedures and policies
3/12/04	FDA Fax regarding Quality of Life analyses in	1/5/04	276	Information Amendment - Clinical
	RESIST 1 study			Request for FDA Comments on Special
				Protocol Assessment for Protocol 1182.12
4/16/04	FDA Fax regarding CMC comments regarding	3/19/04	340	Info Package for Tyne B Pre-NDA Mtg
	upcoming meeting	4/13/04	381	Amendment to Info Package for Type B Pre-
				NDA Mtg.
5/18/04	FDA Fax regarding Pharmacology/Toxicology	2/20/04	318	Request for FDA Feedback on BI plans for
	Comments SN 318			immunotoxicology testing
6/3/04	FDA Fax regarding DSI request for investigator	12/21/04	21-814	Original NDA
6/8/04	FDA Fax – List of Attendees from line 2 2004	7/19/04	005 NO	Pre.NDA meeting Minutes
			200	
	(SVI) (979)			

Description of Reference			
Serial	Number	Jo	Reference
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Description of FDA Correspondence			
Date of FDA	Correspondence		

6/22/04	FDA Fax – CMC questions for TPV Oral Solution face-to-face meeting – June 24, 2004	6/11/04	SN 460	Information Amendment CMC Type A IND Meeting Information Package for Oral Sol.
7/2/04	FDA Fax – Pharmacology/Toxicology comment reqarding SN 470	6/17/04	470	Information Amendment: PharmTox 24- month oral Rat Carcinogenicity Study – Request for FDA Feedback on planned termination of study arm due to mortality rates.
8/12/04	FDA Fax – Microbiology Comments for IND 51,979 SN 534	7/28/04	534	Request for FDA Feedback - Proposal for in vitro study of ARV's
8/12/04	FDA Fax Microbiology Comments for IND 51,979 SN 527	7/22/04	527	Information Amendment: Pharmacology/Toxicology –(U03-3565, U03-3153)
8/30/04	FDA Letter – PRE-NDA CMC Meeting (type B) meeting minutes – April 19, 2004	4/19/04 3/19/04	N/A 314	Pre-NDA Meeting Pre-NDA Background Document
10/08/04	FDA Fax – DDMAC comments regarding the press release for treatment IND 70,629 for TPV expanded access program	9/30/04	SN002	Submission of Press Release
10/08/04	FDA Fax – Treatment IND 70,629 TPV expanded access program	9/7/04	N/A	Original Treatment IND 70,629
10/20/04	FDA Fax - Clinical Request for Pre-submission data - # of deaths for NDA 21-814	N/A	N/A	N/A
10/25/04	FDA Fax – Clinical and Clinical Pharmacology comments for IND 51,979, SN 577	9/10/04	577	Protocol Amendment: Final Draft Protocol 1182.60 (QT Prolongation Study)
11/5/04	FDA Fax – Clinical Comments for latest IB IND 51,979, SN 594	11/1/04	594	Investigators Brochure - Version 8

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Date of FDA	Correspondence		

11/12/04	FDA Fax – Clinical Request for NDA 21-814 for TPV, CRT datasets difficult to review.	12/21/04	21-814	Original NDA
	requested modification of datasets to facilitate review			
11/12/04	FDA Letter- Official minutes – Pre-NDA meeting June 2, 2004	6/2/2004	N/A	Pre-NDA Meeting
		7/19/01	521	Bl's Pre-NDA Meeting Minutes
11/15/04	FDA Letter - Acknowledge receipt of NDA	12/21/04	21-814	Original NDA – Capsules
	(21814, 21822)	12/21/04	21-822	Original NDA – Solution
11/23/04	Email: Clinical Comments in regards to meeting held on November 22, 2004	11/22/04	N/A	FDA Meeting
11/24/04	Email: Statistics Request	12/21/04	N/A	Original NDA
11/30/04	Email: Questions for Medical/Statistics	12/21/04	21-814	Original NDA
	Reviewers			•
12/01/04	Email: TPV Request re: Microbiology	12/21/04	21-814	Original NDA
12/02/04	Email: TPV query 12/2/04 re 1182.12 Study	12/21/04	21-814	Original NDA
	report			
12/02/04	Email: Questions from Medical/Statistics	11/30/04	N/A	N/A
	Reviewers			
12/17/04	FDA Letter - Official Telecon minutes -	12/21/04	21-814	Original NDA
	withdraw of NDA	12/17/04	N/A	Telecon
12/21/04	FDA Fax – Acknowledge receipt of BIPI	12/21/04	21-814,	Original NDA's
	correspondence notifying FDA of withdrawal of	12/20/04	21-822	Withdrawal of NDA's
	NDA's			
2/8/05	FDA FAX – Tentative Date for TPV Advisory	N/A	N/A	N/A
	Committee Meeting			
2/20/05	FDA Fax - Letter informing of upcoming FDA	N/A	N/A	N/A
	advisory committee meeting			

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3/4/05	FDA Acknowledgement of Receipt of 12/20/04	N/A	N/A	N/A
	Withdrawal and 12/21/04 Resubmission. FDA			
	will give Priority Review.			
3/7/05	FDA FAX – TPV 74 day filing letter	12/21/04	21-814	Original NDA
3/24/05	FDA FAX - Pharm/Tox comments for	12/21/04	21-814	Original NDA
	NDA 21-814 for TPV- Comments on behalf of			)
	Dr. Bigger			
4/22/05	ACR - FDA Background Document for	N/A	N/A	N/A
	upcoming 5/19/05 Advisory Committee			
	Meeting.			
5/2/05	FDA fax – Agency backgrounder	N/A	N/A	N/A
	Background document provided by CDER to			
	the Advisory Committee Members for the			
	upcoming 5/19/05 meeting			
6/22/05	NDA Approval Letter	12/21/04	21-814	Original NDA
6/23/05	FDA letter stating that labeling attached to the	6/22/05	21-814	NDA Approval Letter
	approval letter was incorrect			

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<b>Tipranavir</b>

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference	
100	10/27/04	• Response to FDA Request for Information – deaths from studies .12, .48, .17,.1, .4, .5, .51, .52, and .58 along with all 7day faxes of deaths and narratives for each death	FDA Fax	10/20/04	Response to FDA Fax dated October 20, 2004 requesting info on TPV deaths in studies 1182.12, 1182.48, 1182.17 and 1182.58	
002	10/29/04	• Replacement Volume – Module 2 Volume 1.6 – Summary of Clinical Safety	A001	10/21/04	Safety (replacement volume to correct technical error in cross referencing of original 1.6 submitted in original NDA dated October 21, 2004.	
003	11/01/04	<ul> <li>Electronic Submission Update:Corporate Drug Safety Dataset</li> </ul>	Original NDA	10/21/04	Original dataset included in October 21, 2004 Original NDA	
			A001	10/27/04	Updated electronic dataset to include info in Amendment 1	
004	11/03/04	<ul> <li>General Correspondence / Request for FDA Feedback: Proposal for 2 Month Safety Update</li> </ul>	Pre-NDAMeeting	6/2/04	At meeting BI discussed the request for priority review and in preparation, BI will submit proposal for 2 Month Safety Update	
			SN 521	7/15/04	Pre-NDA Meeting Minutes	

	mission Log
	<b>Fipranavir Capsules NDA 21-814 Submission Log</b>
	Capsules ND
	Tipranavir (

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
005	11/12/04	• Response to FDA Request for Information: Updated Information on Tipranavir Deaths	A001 E-mail	10/27/04	Per e-mail request from Ms. Tanima Sinha dated November 5, 2004, requesting additional information for Dr. Andrea James in reference to Amendment 1
900	11/17/04	<ul> <li>Response to FDA Request for Information: Updated Information on Tipranavir Datasets</li> </ul>	FDA Fax	11/12/04	Fax request of Ms. Sinha dated November 12, 2004 per Clinical reviewers' having difficulty reviewing some of the datasets
002	11/18/04	<ul> <li>General Correspondence / Request for Feedback: Agenda Topics to be discussed on the requested meeting</li> </ul>	Agency Contact	11/17/04	Per telephone discussion with Ms. Tanima Sinha dated November 17, 2004, requesting a meeting with BI on November 22, 2004, summary of understanding of issues identified with NDA submission.
800	11/30/04	General Correspondence:  Justification for Priority Review	Telecon	11/24/04	Telecon with FDA 11/24/04: Agreement made that BIPI would submit a short justification for priority Review.
600	12/03/04	General Correspondence: Proposed Electronic Submission Timeline	FDA Meeting Telecon Agency	11/22/04 11/23/04 12/03/04	As agreed during telephone discussion with Drs. Blank and Kaplan, and per discussion with Ms. Sinha, BI is providing a description of all action items agreed between the Division and BI at the meeting

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
			Contact		of November 22 and in communication since that time.
010	12/03/04	<ul> <li>Response to FDA Request for Information: Updated Resistance Dataset</li> </ul>	e-mail	12/01/04	Microbiology Reviewers', Kimberly Struble and Lisa Naeger, comments regarding the Tipranavir BASE-RES Dataset.
011	12/05/04	Response to FDA Request for Information – Updated Information of TPV Datasets: Response to Requests from Nov 22, 2004 meeting	Telefax FDA Meeting	11/12/04	Facsimile request from ms. Tanima Sinha, dated November 12, 2004 with Reviewers' comments regarding the CRT datasets.
012	12/06/04	• Response to FDA Request for Information – Comments regarding 1182.12 study report	e-mail	12/02/04	E-mail request from Ms. Tanima Sinha dated December 2, 2004, on behalf of clinical Reviewer, Dr. James with comments regarding the 1182.12 study report
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		Tipranavir Capsules NDA 21-814 Submission Log	NDA 21-81	Tillano t	ssion Log
Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
013	12/06/04	Response to FDA Request for Information – Proposals to items 5, 8 and 9 from our BDM group. Example of SAS dataset	FDA Meeting E-mail Amend 011	11/22/04 11/23/04 12/05/04	E-mail request from Ms. Tanima Sinha dated November 23, 2004, with Reviewers' comments summarized from our November 22, 2004 meeting
014	12/09/04	Response to FDA Request for Information: Updated Information on Tipranavir Datasets – Final changes from FDA meeting on Nov. 22, 2004 + side requests	FDA Meeting E-mail A 011 Telecon	11/22/04 11/23/04 12/05/04 12/07/04	E-mail request from Ms. Tanima Sinha dated November 23, 2004, with Reviewers' comments summarized from our November 22, 2004 meeting. Reference also made to a teleconference held on Dec ember 7, 2004.
015	12/17/04	• Response to FDA Request for Information – PK reviewer Questions of November 24, 2004 and December 3, 2004	E-mail	11/24/04	E-mail request from Ms. Tanima Sinha on behalf of PK reviewer dated November 24, 2004 and December 23, 2004.
N/A	12/20/04	Request to withdraw New Drug Application	Original NDA	10/21/04	Original New Drug Application
N/A	12/21/04	Resubmission of New Drug Application	Original NDA Withdrawal of NDA	10/21/04	Original New Drug Application Withdrawal of New Drug Application
016	12/29/04	Response to Request for	Telecon	12/20/04	Dr. James requested that BI submit a new

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		Tipranavir Capsules NDA 21-814 Submission Log	NDA 21-81	4 Submi	ssion Log
Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
		Information - Replacement E-Submission	E-mail	12/20/04	electronic submission with all updated datasets submitted since October 22, 2004
017	1/4/05	Response to FDA Request for Information – Immunotoxicity Study	SN 381	2/20/04	BI immunotox proposal BI agreed to conduct study
			Telecon	8/30/04	Summary of Immunotoxicity Study 04R103 would be available for Jan. 4, 2004
			E-mail	9/17/04	
018	1/12/05	• Response to FDA Request for Information – Statistical – RESIST 1 and 2	E-mail	1/7/05	Statistical comments/questions refer to data for RESIST 1 (Study 1182.12) and RESIST 2 (Study 1182.48)
019	1/13/05	Response to FDA Request for Information - Statistical request for cutoff dates/ patient exposures for e-datasests	E-mail	1/11/05	Email request from FDA Requesting a table showing the database lock dates and length of follow-up of patients for each electronic data submission to the NDA

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
020	1/25/05	<ul> <li>Response to FDA Request for Information - In Vitro Virologic Interations Study</li> </ul>	Pre-NDA Mtg	6/2/04	N/A
			SN 534	7/28/04	BI submission requesting feedback on the study design and timing, response to request of SN 534
			E-mail	8/12/04	Request for info on in-vitro virologic interactions study
021	1/25/05	• General Correspondence: December 17, 2004 Meeting Minutes / Request for Change to Official FDA Meeting Minutes	Meeting	12/17/04	Meeting between BIPI and Division
22	2/2/05	• Written Responses to November 17, 2004 e-mail.	E-mail	11/17/04	Request from stat reviewer, Dr. Bhore with comments regarding the CRT datasets.
23	2/3/05	<ul> <li>Background Therapy Optimization Tablets</li> </ul>	E-mail	1/31/05	Request from Dr. Liang in preparation for teleconference February 4, 2005. request for tables reformatting data for patients in Resist 1 and Resist 2.

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		Tipranavir Capsules NDA 21-814 Submission Log	NDA 21-81	4 Submi	ssion Log
Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
24	2/9/05	Rheumatology or Dermatology consults for Trial 1182.22	Email	2/3/05	FDA Requesting pictures electronically for section 16.4 of Report 1182.22
25	2/10/05	Response to FDA Optimization and Listings of 1182.12 and 1182.48 RESIST Studies (datasets)	FDA Letter and Telecon	1/31/05, 2/4/05	FDA questioning background therapy optimzation
26	2/10/05	<ul> <li>Response to FDA request regarding RESIST 1 and RESIST 2 discontinuations</li> </ul>	Email	2/8/05	Email Request from Dr. Bhore and Dr. Baylor – requesting RESIST 1 and RESIST 2 discontinuations
27	2/16/06	Response to Request for Information - Resubmission of Rheumatology and Dermatology Consults –	Email	2/3/05	FDA Requesting pictures electronically for section 16.4 of Report 1182.22
28	2/16/05	<ul> <li>Response to FDA Request for Information: Updated Information on Tipranavir Datasets</li> </ul>	Email	2/2/05	Request from Ms. Monica Zeballos with Clinical, Statistical and PK queries.
29	2/22/05	• 2 Month Safety Update Report	A004 Telecon	11/3/04	Requesting FDA feedback on 2MSUR FDA agreed to the proposal and timing for the submission of the 2MSUR

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
30	2/23/05	• Response to Request for Information - Resistance Datasets	Meetings	6/4/04,	Pre-NDA Meeting Electronic submission demonstration
31	2/25/05	• Response to Request for Information on trials 1182.4, 1182.51 & 1182.22	Email	2/14/05	Query from Dr. Gibbs regarding trials 1182.4, 1182.6, 1182.51 and 1182.22
32	2/28/05	<ul> <li>Response to Request for Information on Datasets</li> </ul>	Email	2/25/05	FDA query (Dr. Bhore) regarding Resist I clinical trials (Studies 1182.12 and 1182.48)
33	2/28/05	• Response to Request for Information – updated ARV datasets for trials 1182.51 and 1182.52	E-mail	2/18/05	Updated ARV datasets for trials 1182.51 and 1182.52
34	3/3/05	Response to FDA Request for	Telecon	8/30/04	Discussion of immunotoxicity program

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
		information – Final Immunotoxicity Report U05-3021	SN 318	2/20/04	Submission describing program
		-	SN 582	9/23/04	Requesting feedback on the adequacy of the protocol design
			A017	3/3/05	Summary of study 04R013 (U05-3021).
35	3/4/05	<ul> <li>Response to FDA Request for information. CRF's for 1182.12 subjects.</li> </ul>	E-mail	3/1/05	E-mail request of March 1, 2005 from Clinical Reviewer, Dr. James with a request for case report forms for subjects in the 1182.12 trial.
36	3/15/05	• Response to FDA Request for	A028	2/22/05	2 Month Safety Update Report
		2MSUR	Telecon	3/9/05	N/A
			NDA filing	3/10/05	NDA Filing communication
37	3/15/05	Response to FDA Request for Information; Adverse Event Information for Naïve Study	E-mail	3/10/05	Ms. Tanima Sinha of behalf of Dr. Melisse Baylor requesting information for the Naïve Study (1182.33)

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
		1182.33			7000 0000
38	3/15/05	• Response to FDA Request:- country De-codes	E-mail	3/14/05	Ms. Tanima Sinha on behalf from Dr. Bhore regarding country codes.
39	3/16/05	Response to FDA Request of March 9, 2005: micro reviewer queries	E-mail	3/9/05	Ms. Tanima Sinha on behalf of Dr. Naeger – comments and clarifications requested for report U00-3184.
40	3/17/05	<ul> <li>Response to FDA Request for Information; Request of March 10, 2005 statistical queries.</li> </ul>	E-mail	3/10/05	Comments and queries regarding Resist I clinical trials 1182.12 and 882.48.
14	3/21/05	<ul> <li>CMC Amendment – updated stability - Change in Proposed Storage Conditions</li> </ul>	N/A	N/A	N/A
42	3/24/05	Response to FDA Request for Information:Laboratory Information for Naïve Study.	E-mail A037	3/10/05	e-mail request from Ms. Tanima Sinha on behalf of Dr., Baylor Adverse Event information for Naïve study
43	3/24/05	<ul> <li>Advisory committee meeting package – DRAFT background document for 4/4/05 meeting</li> </ul>	Telecon	3/21/05	Discussed proposed agenda for ADVAC meeting.

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Ssion Log  Description of Reference	N/A	2 Month Safety Update Response to Request for Information - Resistance Datasets	CMC Amendment justifying additional excipient	Pharmacology Toxicology Query on Amendment 044	Micro Requests from Lisa Naeger on phenotypic data.	PK request – revised labeling
4 Submi Date of Reference	N/A	2/22//05	3/24/05	3/30/05	3/23/05	3/16/05
NDA 21-81 Reference	N/A	A029 A030	A044	E-mail	E-mail & Telecon	Telecon
Tipranavir Capsules NDA 21-814 Submission Log  Description of Submission Reference Date of Reference	CMC Amendment - Additional data to justify the specification for an additional excipient	Labeling Update and Resistance Amendment -Tipranavir Analysis of Emergent Resistance/Revised Labeling	Response to FDA Request for Information	Dog Bioavailability Study	<ul> <li>Response to FDA Request for Information – Microbiology – phenotypic data</li> </ul>	Revised Labeling – revised labeling reflecting pk responses
Date of Submission	3/24/05	3/25/05	3/31/05		3/31/05	4/4/2005
Amendment	44	45	46		47	48

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Amendment	Date of	Description of Submission Reference Date of Description	Reference	Date of	Description of Reference
26	4/15/05	Revised Labeling Table 1 (Drug Interactions) Correction to Amendment 048	A048	4/1/05	Revised labeling submitted April 1, 2005, correction to table 1
57	4/19/05	General Correspondence: Tipranavir Advisory Committee Briefing Document	Telecon A053	4/11/05	Teleconference regarding the Resistance Section of the AVDAC Briefing Document Request for FDA Feedback on Resistance Section of AVDAC Briefing Document
28	4/19/05	Response to FDA Request for Information – efficacy analysis from the RESIST program	E-mail Telecon	4/13/05	Dr. Bhore's tipranavir efficacy analysis from the RESIST program Discussed potential reasons for differences in the outcome of Dr. Bhores analysis versus BIPI's.
59	4/19/05	Response to FDA Request for Information – patient disposition	E-mail	4/7/05	Statistical queries from Dr. Bhore on patient disposition
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		Lipranavir Capsules NDA 21-814 Submission Log	NDA 21-81	4 Submi	ssion Log
Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
09	4/19/05	Response to FDA Request for	A49	4/6/05	Reference Amendment 49 on April 6, 2005
		Information – protocol deviation datasets	Telephone Discussion/e-mail	4/18/05	and telephone discussion as well as email on April 18, 2005 asking for new protocol deviation datasets
61	4/21/05	Response to FDA Request 74 Day Filing Letter	Letter	3/7/05	Reference to 74 Day Filing Letter from March 7, 2005
62	4/28/05	Response to FDA Request for Information – OBR Definition	E-mail	4/27/05	Statistical Query - OBR Definition
63	4/29/05	Response to FDA Request for Information – datasets on protocol violations	E-mail	4/29/05	Dr. Andrea James' request that BI direct her to the raw or analysis datasets that will provide information on protocol violations
64	4/29/05	Response to FDA Request for Information – location of tables for the classification of toxicity grading for studies .4, .12, .48, .51 and .52	E-mail	4/29/05	On behalf of Medical reviewer requesting the location of tables for the classification of toxicity grading for studies 1182.4, .12, .48, .51, and .52.
65	5/2/05	Background Documentation for May	A057	4/19/05	TPV Advisory Committee Briefing

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
		4, 2005 Teleconference AVDAC Presentation slides			Document
			Telecon	4/27/05	Current version of AVDAC presentation
99	2/2/02	Response to FDA Request for Information- specific resist patient lab and AES and patient disposition	E-mail	4/29/05	Query from Dr. James regarding specific individual resist patient lab and clinical, AES and patient disposition
29	5/10/05	Response to FDA Request for Information – lipodystrophy data	E-mail	5/10/05	Query from Dr. James regarding data for lipodystrophy
89	5/10/05	Background Information for May 11, 2004 telecon – updated slides for	A057	4/19/05	TPV Advisory Committee Briefing Document
		resistance presentation	A065	5/2/05	Background Document for AVDAC
69	5/11/05	Response to FDA Request for Information – In vitro activity assessments	E-mail	4/4/05	Request for In vitro combination activity assessments
70	5/13/05	Response to FDA Request for Information – response to efficacy	E-mail	4/13/05	Dr. Bhore's tipranvir efficacy analysis from the RESIST program
		aliatysis for NESIST T aliu NESIST 2	Telecon	4/14/05	Discussed potential reasons for differences

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
			A058	4/19/05	in the outcome of Dr. Bhores analysis versus BIPI's Response to FDA Request for Information – efficacy analysis from the RESIST program
			E-mail	5/4/05	E-mail query on behalf of Dr. Bhore regarding efficacy analyses of RESIST 1 and RESIST 2
71	5/13/05	Response to FDA Request for Information – Sample of letter to Investigators re: Drug Metabolism and Interaction	E-mail	3/16/05	Expanded access program Investigators Brochure –EAP Investigators brochure – FDA requests BIPI inform investigators of information on Drug Metabolism and Interaction.
			telecon	4/27/05 &5/24/05	BIPI agreement to inform investigators

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Amendment	Date of Submission	Description of Submission	Reference	Date of Reference	Description of Reference
72	2/16/05	General Correspondence - Additional	A057	4/19/05	ADVAC Briefing Document
		Silues for ADVAC	N/A	4/22/05	FDA's ADVAC Briefing Document
			Telecon	4/27/05	Current version of ADVAC presentation
			A065	5/2/05	Background Documentation for May 4, 2005 Teleconference ADVAC Presentation Slides
			Telecon	5/11/05	Informing FDA BIPI will submit additional slides for ADVAC meeting
73	5/24/05	Response to FDA Request for Information – response to Pharm, Tox	Telefax	5/13/05	FDA Questions, Pharmacology/Toxicology/CMC
74	5/27/05	Response to FDA Request for Information – Response to CMC questions of 5/13/05	Telefax	5/13/05	CMC Questions
75	9/3/02	Response to FDA Request – Revised Labeling	Telecon	50/8/9	Revised labeling to support the scheduled teleconference between the Division and BI on June 8, 2005.
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76	9/01/09	General Correspondence – Post Marketing Commitments – Clinical and Toxicology	Telecon	9/1/9	Ms. Tanima Sinha requested that BI provide a proposal for post-marketing commitments
7.1	6/10/05	CMC – Amendment to a Pending Application – Updated specifications for drug substance, per FDA comments.	E-mail	9/1/05	FDA comments on the specifications for tipranavir drug substance and capsules.
78	9/10/02	Response to FDA Information Request - Environmental Analysis query of 6/7/05	E-mail	90/L/9	FDA's Information Request regarding Environmental Analysis.
79	6/17/05	CMC – Amendment to a Pending Application: Updated specification for drug product dissolution per FDA and BIPI agreement.	A077	6/10/05	BI agreement to FDA's proposed acceptance criterion for drug product specification and drug product dissolution
08	6/21/05	Post marketing commitments	Telecon	6/1/05	FDA request for post-marketing commitments proposal

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81 6/21/05 General Correspondence – Revised A075 Labeling which incorporates all changes agreed to by members Division and BIPI to date.	Description of Submission Reference	Date of Reference	Description of Reference
		9/8/9	Labeling Revision
	ncorporates all by members I to date.	Multiple	Multiple communications via teleconference and e-mail since 6/3/05 submission.

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Description of Reference		
Serial No.	Jo	Reference
Date of	Reference	
NAME OF	CONTACT	PERSON
REASON FOR CONTACT		
DATE		

6/22/05	Amended Approval Letter, Capsules	Ms. Tanima Sinha	NA	NA	NA
		Project Manager			
	Final capsule approval letter	Ms. Tanima Sinha	NA	NA	NA
6/22/05		Project Manager			
6/22/05	Final medical reviewer query; final	Ms. Tanima Sinha	NA	NA	NA
	review of bottle label; final label	Project Manager			
	revisions				
6/16/05	FDA comments from DSRCS	Ms. Tanima Sinha	NA	NA	NA
	regarding the PPI for TPV	Project Manager			
6/10/05	Status of Clin Stat commentsfor	Ms. Tanima Sinha	NA	NA	NA
	upcoming CMC telecom on 6/10/05	Project Manager			
	CMC telecon				
6/10/05	Final Report of Immunotoxicity	Ms. Tanima Sinha	1/4/05	909	IA/Pharmacology/Toxicology
	Study	Project Manager			Study summary of unaudited draft res
9/1/02	Request for Revised Labeling	Ms. Tanima Sinha	NA	NA	NA
	Request for post-marketing	Project Manager			
	commitments proposal				

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NAME OF	CONTACT	PERSON
REASON FOR CONTACT		
DATE		

	AVDAC	DAVDP, AVDAC	4/19/05	A057	Response to FDA Request for
					Information – Draft Advisory Committee Briefing document
			5/2/05	A065	AVDAC slides
			4/22/05	NA	FDA AVDAC Briefing Document
			Multiple	ACR	Meetings, telecons, to address Upcoming AVDAC
FDA stat response on primary efficacy results of the resist trials/amendment 058 for telecon – 11 May 2005	n primary te resist 88 for telecon –	Ms. Tanima Sinha Project Manager	4/19/05	A057	Response to FDA Request for Information – Draft Advisory Committee Briefing document:
s for	n team and order ADVAC meeting	Ms. Anuja Patel, MPH Health Science Administrator	4/19/05	A057	General Correspondence:TPV Advisory Committee Meeting Briefing document (DRAFT)

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	NA	N/A	General Correspondence: Advisory Committee Briefing Document - DRAFT	NA	Response to FDA Request For Information: Comments Regarding 1182.12 study report	Response to FDA Request For Information: datasets that will provide info on protocol violations	Response to FDA Request for Information – Efficacy Analysis from RESIST program
	NA	N/A	057	NA	A012	A063	A058
	NA	N/A	4/19/05	NA	12/6/04	4/29/05	4/19/05
	Ms. Tanima Sinha Project Manager	Elizabeth Thompson Project Manager, DAVDP	Ms. Tanima Sinha Project Manager	Ms. Anuja Patel, MPH Health Science Administrator	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager
	FDA query on Lipodystrophy information in datasets from Dr. James	FDA request to reschedule telecon to 6 May 2005 at 9:00am	Request for FDA feedback on BI AVDAC presentation/Request for copy of FDA's AVDAC presentation	Request for List of investigator and sub-investigators in TPV clinical program	FDA query – tables for the classification of toxicity grading for studies 0040, 0012, 0048, 0051 and 0052	FDA query from Dr. James regarding raw or analysis datasets/CMC and pharm\tox comments should come next week	FDA request for emailed copy of amendment 058
	5/10/05	5/3/05	5/2/05	4/29/05-5/1/05	4/29/05	4/29/05	4/29/05

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		1		τ	1	r	
	NA	Response to FDA Request for New Protocol deviation datasets	General Correspondence: Advisory Committee Briefing Document - DRAFT	NA	NA	General Correspondence: tipranavir Advisory Committee Meeting Briefing Document - DRAFT	General Correspondence: tipranavir Advisory Committee Meeting Briefing Document - DRAFT
Reference	NA	4/19/05	4/12/05	NA	NA	A057	A057
	NA	090	A057	NA	NA	4/19/05	4/19/05
PERSON	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Anuja Patel, MPH Health Science	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Anuja Patel, MPH Health Science Administrator
	FDA query from Dr. James regarding specific individual RESIST PT lab (AST, ALT, BILI) and clinical (specifically AES, and disposition)	FDA stat query from Dr. Bhore – A060 Protocol deviations dataset	May 19 AVDAC meeting – contact information for industry rep	Discussion of CARC studies for traditional approval	Inquiring when BIPI will receive CMC and non clinical queries for TPV	FDA receipt of AO57 Briefing Document	FDA receipt of 19 April AVDAC Briefing DocumenVFDA request for additional CD
	4/29/05	4/27/05	4/27/05	4/26/05	4/25/05	4/20/05	4/20/05 & 4/25/05

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4/19/05	AO57 AVDAC	Ms. Tanima Sinha	4/19/05	A057	General Correspondence:
	Briefing Document	Project Manager			tipranavir Advisory
					Committee Meeting Briefing
1000					Document - DKAF1
4/19/05	AVDAC briefing document –	Ms. Tanima Sinha	4/19/05	A057	General Correspondence:
	solution NDA # in header	Project Manager			tipranavir Advisory
			-		Committee Meeting Briefing
					Document - DRAFT
4/19/05	FDA request to cancel weekly	Ms. Tanima Sinha	NA	NA	NA
	clin/stat teleconference on 20 April,	Project Manager			
	2005				
4/18/05	BI response to reminder that AVDAC Ms. Anuja Patel,	Ms. Anuja Patel,	NA	NA	NA
	backgrounder due 19 April	MPH			
		Health Science			
		Administrator			
4/18/05	FDA TPV request for patient	Ms. Tanima Sinha	4/6/05	A049	Response to FDA Request
	numbers from NDA A049 (Protocol	Project Manager			For Information: response
	Deviations) table 4 (PTS with				to protocol deviation queries
	relevant protocol deviations for				•
	Resist 1 and Resist 2				

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		TOOM T			
4/15/05	TPV Peds query and DDMAC Interaction	Ms. Tanima Sinha Project Manager Dr. Rosemary Johann-Liang Medical Team Leader Dr. Melisse Baylor Medical Reviewer	4/4/05	N/A	Meeting with Division And BIPI
4/14/05	FDA stat query A052 response to April 6, 2005 stats query (genotype sensitivity score)	Ms. Tanima Sinha Project Manager	4/12/05	A052	Response to FDA Request For Information: statistical Queries from Ms. Zeballas of 4/7/05 via e-mail
4/13/05	FDA comments regarding AC Backgrounder for TPV-STATS efficacy comments	Ms. Tanima Sinha Project Manager	4/19/05	A057	General Correspondence: Tipranavir Advisory Committee Meeting Briefing Document - DRAFT
4/13/05	FDA request for telecon TPV RESIST efficacy results and outcome	Ms. Tanima Sinha Project Manager	NA	NA	NA
4/13/05	FDA follow up issues – STAT comment on April 7, 2005 query – Patient disposition	Ms. Tanima Sinha Project Manager	4/7/05	NA	Stat comments on April 7, 2005 query
4/13/05	FDA microbiology comments regarding AC briefing document	Ms. Tanima Sinha Project Manager	4/19/05	A057	General Correspondence Tipranavir Advisory Committee Briefing document - DRAFT

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General Correspondence Tipranavir Advisory Committee Briefing document	V.	NA	NA	DRAFT ADVAC Background document for 4/4/05 meeting	meeting with FDA to review background document
A043	Υ <sub>N</sub>	NA	NA	A043	meeting
3/24/05	NA A	NA	NA	3/34/05	4/4/05
Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	
FDA comments regarding AC backgrounder for TPV – Clinical AE grading	Follow up on  Request for feedback on timing and need for updated disposition dataset  AVDAC preparation — Request for feedback on resistance package and FDA's safety presentation  Written response that final CARC studies not needed for TA	FDA weekly clin/stat telecon regarding TPV 27 April 2005	FDA query – comments on pediatric study report	FDA comments on AC backgrounder 4/11/05	
4/13/05	4/13/05	4/11/05	4/11/05	4/11/05	

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		PERSON		Reference	
4/4/05	Various questions/comments at side	Ms. Tanima Sinha	3/24/05	A043	Background Document for
	meeting during 4 April 05	Project Manager			4/4/05 meeting
	BI_DAVDP meeting, importantly				
	including the topics of Traditional		4/4/05	NA	April 4, 2005 Meeting
	Approval, Monthly Safety Reports,				
	AVDAC Rehearsing, Meeting				
	Participants				
4/1/05	FDA request – TPV SAE fatal	Ms. Tanima Sinha	NA	NA	NA
		Project Manager			
3/30/25	FDA TPV Query from Dr. Gibbs -	Ms. Tanima Sinha	NA	NA	NA
	Trials 1182.4, 1182.6 and	Project Manager			
	1182.51/Queries on 1182.22	,			
3/23/05	Micro Requests RE varied>fold	Ms. Tanima Sinha	NA	NA	NA
	change values in pheno data/missing	Project Manager			
	pheno data				
3/16/05	FDA comments regarding EAP and	Ms. Tanima Sinha	3/16/05	e-mail	FDA comments regarding EAP
	the IB for TPV – updates needed –	Project Manager			and the IB for TPV - updates
	corrected e-mail from original 3/16				needed
	query.				
3/16/05	FDA stats query – data coding	Ms. Tanima Sinha	NA	NA	NA
	problems	Project Manager			
3/14/05	STAT queries on TPV NDA 21-814	Ms. Tanima Sinha	NA	NA	NA
	from Dr. Bhore – data for RESIST 2	Project Manager			
	trial (study 1182.48)				
3/10/05	TPV query from Dr. James - Dataset	Ms. Tanima Sinha	NA	NA	NA
	Discrepancies	Project Manager			

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NA		NA		NA			NA		NA		NA		NA		NA		NA		NA	,	NA			
NA		NA		NA			NA		NA		NA		NA		NA		NA		NA		NA			
Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager		Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager		
Stat queries on TPV resist trials -	UBK	TPV ACR – AE Information for	ivaive study/updated tabei	Microbiology	comments/information/clarification	request for TPV	FDA TPV NDA 21-814 queries – 74	day filing review issues letter	TPV NDA 21-814 queries regarding	pharm/tox or chemistry	Request for Clarification of Variables	referring to ITT Population	FDA canceling weekly clin/stat	telecon-Wednesday, 2/23	FDA comments from Dr. Baylor for	TPV Studies 1182.51 and 1182.52	request from Dr.Zhang-PK	Reviewer	Propose Cancellation of this week's	teleconference	FDA TPV Query from Dr. Gibbs -	Trials 1182.4,1182.6 and	1182.51/Queries on 1182.22	
3/10/05		3/10/05		3/9/05			3/4/05		2/25/05		2/25/05		2/18/05		2/18/05		2/16/05		2/15/05		2/14/05			

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2/11/05	Study Reports – Phase III studies 1182.12 & 1182.48 – FDA cannot locate in NDA.	Ms. Tanima Sinha Project Manager	12/21/04	NA	Original NDA
2/9/05	FDA information requests  ? 1182.51: provide measuring code in DESTERM. ? 1182.22 – Requesting temperature measurements taken during trial	Virginia Behr Chief Project Manager, DAVDP	NA	NA A	NA
2/9/05	Several topics discussed- 2MSUR,IND AR, Filing Date, AVDAC prep, resistance, site inspections, responses to requests for information	Ms. Tanima Sinha Project Manager	NA	NA	NA
2/9/05	TPV NDA – 21-814 – Participants at 4 Feb. Telecon	Ms. Tanima Sinha Project Manager	NA	NA	NA
2/9/05	TPV NDA-21-814 AVDAC tentative meeting agenda	Ms. Tanima Sinha Project Manager	NA	NA	NA
2/8/05	FDA Information Request from Dr. Bhore and Dr. Baylor – RESIST 1 and RESIST 2- regarding discontinuations/additions/switches of antiretrovirals on study.	Ms. Tanima Sinha Project Manager	NA	NA	NA
2/8/05	FDA Request for Info for CRF for PT   Ms. Tanima Sinha #2052-Trial 1182.22   Project Manager	Ms. Tanima Sinha Project Manager	NA	NA	NA

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2/3/05 Clarification of format for amendment 023/Agreement to clarify request for dataset (Question 11) in FDA's Feb 2, 2005 email request at teleconference on Feb 4, 2005  2/3/05-2/4/05 Voicemail regarding NDA 21-814 for Janet Rowe NA an Audit FDA email requesting location of Virginia Behr 12/21/04 information for the 1182.22 study Manager, DAVDP Correction to FDA's Feb. 2, 2005 Ms. Tanima Sinha 2/2/05 email List of Queries Project Manager Pariday at 9:30/clin/stat queries - OBR, ARV changes, clarification of Manager responders  2/1/05 FDA teleconference rescheduled for Regulatory Project OBR, ARV changes, clarification of Manager responders  2/1/05 Teleconference Details - FDA Foiect Manager Project Manager Project Manager Project Manager Project Manager Project Manager Project Manager DAVDP Project Manager Project Manager Project Manager Project Manager Project Manager DAVDP Project Manager Project Manager DAVDP Project Manager Proje			I LINDOIN		יייויייייייי	
Clarification of format for amendment 023/Agreement to clarify request for dataset (Question 11) in FDA's Feb 2, 2005 email request at teleconference on Feb 4, 2005  2/4/05 Voicemail regarding NDA 21-814 for Janet Rowe an Audit information for the 1182.22 study Chief Project Manager, DAVDP Correction to FDA's Feb. 2, 2005  FDA email List of Queries Project Manager FDA teleconference rescheduled for Monica Zeballos Friday at 9:30/clin/stat queries - Regulatory Project OBR, ARV changes, clarification of Manager responders Teleconference Details - FDA Elizabeth proposing time change Project Manager Confirm meeting arrangements for Project Manager Project Manager Confirm meeting arrangements for Project Manager Jan. 28, 2005 teleconference Project Manager Proj						
request for dataset (Question 11) in FDA's Feb 2, 2005 email request at teleconference on Feb 4, 2005  an Audit FDA email requesting location of information for the 1182.22 study Correction to FDA's Feb. 2, 2005  FDA teleconference rescheduled for FDA teleconference rescheduled for FDA teleconference Details – FDA  Teleconference Details – FDA  Call placed informing 7-Day Safety Report would be faxed.  Confirm meeting arrangements for Project Manager  Broject Manager  Confirm meeting arrangements for Project Manager  Report would be faxed.  Project Manager  Report Wanager  Project Manager  Report would be faxed.  Project Manager  Report Wanager  Project Manager  Project Manager  Report would be faxed.  Project Manager	2/4/05	Clarification of format for	Janet Rowe Program Specialist	2/3/05	023	Response to FDA Request
FDA's Feb 2, 2005 email request at teleconference on Feb 4, 2005  Voicemail regarding NDA 21-814 for Janet Rowe an Audit FDA email requesting location of information for the 1182.22 study Manager, DAVDP Correction to FDA's Feb. 2, 2005  FDA teleconference rescheduled for Friday at 9:30/clin/stat queries - OBR, ARV changes, clarification of Fizabeth proposing time change  Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed.  Confirm meeting arrangements for Project Manager Confirm meeting arrangements for Project Manager Project Ma		request for dataset (Question 11) in	i iogianii opedansi			data for patients in the
teleconference on Feb 4, 2005  Voicemail regarding NDA 21-814 for Janet Rowe an Audit  FDA email requesting location of virginia Behr information for the 1182.22 study  Correction to FDA's Feb. 2, 2005  Correction to FDA's Feb. 2, 2005  FDA teleconference rescheduled for Manager responders  Fiday at 9:30/clin/stat queries — Regulatory Project Manager responders  Teleconference Details — FDA  Friday at 9:30/clin/stat queries — Regulatory Project Manager responders  Teleconference Details — FDA  Friday at 9:30/clin/stat queries — Regulatory Project Manager responders  Teleconference Details — FDA  Friday at 9:30/clin/stat queries — Regulatory Project Manager proposing time change — Thompson Project Manager — DAVDP  Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed — Ms. Tanima Sinha Jan. 28, 2005 teleconference — Project Manager — Project Man		FDA's Feb 2, 2005 email request at				RESIST 1 and RESIST 2
2/4/05 Voicemail regarding NDA 21-814 for Janet Rowe an Audit FDA email requesting location of information for the 1182.22 study Correction to FDA's Feb. 2, 2005 Friday at 9:30/clin/stat queries — Regulatory Project Manager FDA teleconference rescheduled for Monica Zeballos Friday at 9:30/clin/stat queries — Regulatory Project OBR, ARV changes, clarification of Manager responders Teleconference Details — FDA proposing time change Call placed informing 7-Day Safety Report would be faxed. Call placed informing 7-Day Safety Report would be faxed. Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference Project Manager Project Manager		teleconference on Feb 4, 2005				Programs Background Therapy Optimization tables
an Audit  FDA email requesting location of information for the 1182.22 study Correction to FDA's Feb. 2, 2005  Correction to FDA's Feb. 2, 2005  Correction to FDA's Feb. 2, 2005  Friday at 9:30/clin/stat queries - Regulatory Project Manager  Responders  Teleconference Details - FDA  Project Manager  Teleconference Details - FDA  Project Manager  Thompson  Project Manager  Thompson  Project Manager  Thompson  Project Manager  Thompson  Project Manager  As Tanima Sinha  Report would be faxed.  Confirm meeting arrangements for Project Manager  Confirm meeting arrangements for Project Manager	2/3/05-2/4/05	iil regardin	Janet Rowe	NA	NA	NA
FDA email requesting location of information for the 1182.22 study information for the 1182.22 study Chief Project Manager, DAVDP Manager, DAVDP Manager, DAVDP Project Manager FDA teleconference rescheduled for Monica Zeballos Friday at 9:30/clin/stat queries – Regulatory Project OBR, ARV changes, clarification of responders Teleconference Details – FDA Elizabeth proposing time change Project Manager, DAVDP Call placed informing 7-Day Safety Ms. Tanima Sinha Beport would be faxed.  Confirm meeting arrangements for Project Manager Projec		an Audit	Program Specialist			
information for the 1182.22 study  Correction to FDA's Feb. 2, 2005  Correction to FDA's Feb. 2, 2005  Foreign Manager  FDA teleconference rescheduled for Monica Zeballos Friday at 9:30/clin/stat queries — Regulatory Project OBR, ARV changes, clarification of Manager responders  Teleconference Details — FDA  Project Manager, DAVDP  Call placed informing 7-Day Safety Report would be faxed.  Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference Project Manager Project Manager Ms. Tanima Sinha Project Manager	2/3/05	FDA email requesting location of	Virginia Behr	12/21/04	NA	Original NDA
Correction to FDA's Feb. 2, 2005  Roadil List of Queries  FDA teleconference rescheduled for Friday at 9:30/clin/stat queries — Regulatory Project OBR, ARV changes, clarification of Fizabeth proposing time change — Thompson Project Manager, DAVDP  Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed.  Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference Prize Manager Project Manag		information for the 1182.22 study	Chief Project			
Correction to FDA's Feb. 2, 2005  Ms. Tanima Sinha email List of Queries  FDA teleconference rescheduled for Friday at 9:30/clin/stat queries – Regulatory Project OBR, ARV changes, clarification of responders  Teleconference Details – FDA Project Manager  Teleconference Details – FDA Project Manager  Call placed informing 7-Day Safety Report would be faxed.  Confirm meeting arrangements for Jan. 28, 2005 teleconference Project Manager Project Manager Project Manager Project Manager Project Manager			Manager, DAVDP			
EDA teleconference rescheduled for Monica Zeballos Friday at 9:30/clin/stat queries – Regulatory Project OBR, ARV changes, clarification of Manager responders Teleconference Details – FDA Project Manager, DAVDP Call placed informing 7-Day Safety Report would be faxed. Confirm meeting arrangements for Project Manager Jan. 28, 2005 teleconference Project Manager Project Manager Project Manager Project Manager Project Manager	2/3/05	Correction to FDA's Feb. 2, 2005	Ms. Tanima Sinha	2/2/05	FDA e-mail	Clin/Stat Queries - OBR, ARV
FDA teleconference rescheduled for Friday at 9:30/clin/stat queries – Regulatory Project OBR, ARV changes, clarification of Frizabeth proposing time change Project Manager, DAVDP Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed.  Confirm meeting arrangements for Project Manager Jan. 28, 2005 teleconference Project Manager P		email List of Queries	Project Manager			Changes, clarification of
FDA teleconference rescheduled for Monica Zeballos Friday at 9:30/clin/stat queries – Regulatory Project OBR, ARV changes, clarification of Manager responders Teleconference Details – FDA Project Manager, DAVDP Call placed informing 7-Day Safety Report would be faxed. Confirm meeting arrangements for Project Manager Jan. 28, 2005 teleconference Project Manager Project Manager Project Manager Project Manager Project Manager						responders
Friday at 9:30/clin/stat queries – Regulatory Project OBR, ARV changes, clarification of Manager responders Teleconference Details – FDA proposing time change Friday at 9:30/clin/stat queries – Regulatory Project Manager Call placed informing 7-Day Safety Report would be faxed. Sconfirm meeting arrangements for Project Manager Jan. 28, 2005 teleconference Project Manager Project Manager Project Manager	2/2/05	FDA teleconference rescheduled for	Monica Zeballos	N/A	N/A	N/A
OBR, ARV changes, clarification of responders Teleconference Details – FDA Elizabeth proposing time change Project Manager, DAVDP Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed. Confirm meeting arrangements for Project Manager Jan. 28, 2005 teleconference Project Manager Project Manager Project Manager Project Manager Project Manager		Friday at 9:30/clin/stat queries –	Regulatory Project			
responders  Teleconference Details – FDA  Elizabeth  Thompson  Project Manager,  DAVDP  Call placed informing 7-Day Safety  Report would be faxed.  Confirm meeting arrangements for Jan. 28, 2005 teleconference  Project Manager  Ms. Tanima Sinha  Project Manager		OBR, ARV changes, clarification of	Manager			
Teleconference Details – FDA Elizabeth proposing time change Thompson Project Manager, DAVDP Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed. Project Manager Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference Project Manager		responders			•	
Project Manager,  Project Manager,  DAVDP  Call placed informing 7-Day Safety  Report would be faxed.  Confirm meeting arrangements for Janima Sinha  Project Manager  Confirm 28, 2005 teleconference  Project Manager	2/1/05	Teleconference Details - FDA	Elizabeth	N/A	N/A	N/A
Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed. Project Manager Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference Project Manager		proposing time change	Thompson			
Call placed informing 7-Day Safety Ms. Tanima Sinha Report would be faxed.  Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference			Project Manager, DAVDP			
Report would be faxed. Project Manager Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference Project Manager	1/28/05	Call placed informing 7-Day Safety	Ms. Tanima Sinha	N/A	N/A	2005-BP-00649RA(0)
Confirm meeting arrangements for Ms. Tanima Sinha Jan. 28, 2005 teleconference		Report would be faxed.	Project Manager			
	1/27/05	Confirm meeting arrangements for	Ms. Tanima Sinha	N/A	N/A	N/A
		Jan. 28, 2005 teleconference	Project Manager			

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04 e-mail Request from Dr. Bhore – Clarification on BI responses to		ole multiple all electronic submissions	N/A N/A		N/A 2004-FF-00714FF		04 21-814 Original NDA				AA014Response to FDA Request	For Information – updated Information on datasets	N/A N/A	04 A011 N/A
11/14/04		multiple	N/A		N/A		12/21/04				12/9/04		N/A	12/11/04
Ms. Tanima Sinha Project Manager		Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha	Project Manager			Ms. Tanima Sinha	Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha
Arrangements for Jan. 28, 2005 teleconference/Dr. Bhore seeking clarification on BI responses to her	recent comments	Request for Database lock dates and length of patient follow up for each esubmission	Request for Weekly ClinStat	teleconference between Division and BI	Call placed informing 7-Day Safety	Report would be faxed.	Follow up on adequacy of e-	submission datasets	Request for telecon to discuss NDA	options on Friday, December 17, 2004	TPV NDA 21-814 amendment 014	and DSI Information	FDA request for clarification TPV NDA 21-814 – full master files of 1182.12/.48 and .52	FDA teleconference confirmation – 7
1/27/05		1/11/05	1/10/05		12/15/04		12/13/04				12/10/04		12/6/04	12/6/04

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#3 stat query for efficacy data on	Ms. Tanima Sinha	N/A	N/A	N/A
TPV for RESIST 1 and RESIST 2 studies	Project Manager		4	
Materials for 7 December telecon- Biometrics & Data Management	Ms. Tanima Sinha Project Manager	Telefax	11/12/04	Fax request from Ms. Sinha with Reviewers' comments regarding the CRT datasets
		12/05/04	A011	Response to FDA Request for Information of TPV Datasets:
Timeline for e-sub datasets	Ms. Tanima Sinha Project Manager	multiple	multiple	Electronic submissions
FDA request issues of microbiology	Ms. Tanima Sinha	N/A	N/A	N/A
FDA Submission timing/Medical server major outage	Froject Manager Ms. Tanima SinhaProject Manager	N/A	N/A	N/A
FDA Email Re: Datasets submission for TPV – If the proposed Dec 3, 04 submission of datasets is not sent to FDA by COB December 6, 04, it will affect the review of the NDA.	Ms. Tanima Sinha Project Manager	11/12/04	FDA email	Request for datasets

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12/02/04	FDA Request TPV Clarifications Datasets	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
	confirming that the real raw datasets				
	correspond exactly to the CRF and				
	the submitted Clinical Study Reports				
	and Integrated Summary of Safety				
12/01/04	TPV Updated Datasets - Due to	Ms. Tanima Sinha	11/12/04	FDA	Request for Datasets
-	outage on storage area network, BIPI	Project Manager		Email	
	cannot commit to the submission of				
	updated datasets by 12/3/04				
11/30/04	TPV Telecon September 30 <sup>th</sup> , 2004		7/28/04	SN 533	General Correpondence - Request
	regarding microbiology comments				for FDA Feedback - Resistance
	for SN 533 and the resistance				Template
	template				
11/30/04	FDA Request for contact Info	Anuja Patel	N/A	N/A	N/A
	March 10 AVDAC Meeting	Executive			
		Secretary AVDAC			
11/24/04	Clin pharm questions: oral solution	Ms. Tanima Sinha	N/A	N/A	N/A
	request for .CTL file, clarification of	Project Manager			
	clin pharm study report				
11/24/04	Called to Set-up site audits	Tony El-Hage	N/A	N/A	N/A
		DSI			

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11/19/04	Telecon on datasets	DAVDP	3/17/04	346	Pre-NDA meeting package
			6/18/04	471	E-sub Plan Update
			7/12/04	ACR	Telecon to discuss E-sub & Adequacy of CDISC 3.0
			9/13,16, 20/04	ACR	Patient profiler / Test DVD
11/17/04	Comments from the clinical review team –Dataset Queries	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
11/17/04	Study report query related to demographic datasets	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA

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11/17/04	FDA #1 Stat Queries for TPV NDA 21-814 – raw datasets and patient disposition file DS. Dr. Bhore needs more clarification.	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
11/12/04	FDA request for Actual Label for Packaging	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
11/5/04	FDA memo from Dr. James regarding NDA for TPV – Request for full narratives, discrepancies in fatality numbers, request for CRFs for fatal cases	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
11/5/04	Cross Referencing for SCS – deaths submission query	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
10/28/04	for SCS – notified the cross- summary of not resolve Il send updated A Amendment.	Ms. Tanima Sinha Project Manager	12/21/04	21-814	Original NDA
10/28/04	NDA Amendment 001 – Submission Clarification	Ms. Tanima Sinha Project Manager	10/28/04	A001	Response to FDA Request for Information on TPV deaths in studies 1182.12, 1182.48, 1182.17 and 1182.58
10/25/04	Call placed informing 7-Day Safety Report would be faxed.	Ms. Tanima Sinha Project Manager	10/28/04	593	IND Safety Reports (10/16/04-10/30/04) Case IW-2004-BP-08523B0

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N/A	N/A	N/A	Pharmacology/Toxicology Request for Feedback on Immunotoxicity Study Protocol	Pre-NDA meeting package	E-sub Plan Update	Telecon to discuss E-sub & Adequacy of CDISC 3.0	Patient profiler / Test DVD	GG E-sub demo agenda, Reviewer's guide, e-sub structure
N/A	N/A	N/A	582	346	471	ACR	ACR	583
N/A	N/A	N/A	9/23/04	3/17/04	6/18/04	7/12/04	20/04	9/27/04
Tom Selnekovic Electronic Document Room Project Manager	Tom Selnekovic Electronic Document Room	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	DAVDP IT				
TPV "test DVD" successfully loaded by FDA	Request to load "test esub DVD"	Telecon – Investigators Brochure	Immunotox Study Proposal – SN 582	Electronic submission demonstration	<i>:</i>			
10/18/04	10/12/04 – 10/ 13/04	10/7/04	10/7/04	10/5/04				

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10/2/04	Meeting to discuss pediatric written	DAVDP	11/15/02	186	Pediatric Proposal
			1/22/03	N/A	Pediatric written request
			7/23/03	242	Request for changes to written request
9/24 &9/27/04	Early Submission of Module 4 –	Ms. Tanima Sinha	N/A	N/A	N/A
	notifying FDA that BIPI Will be submitting Module 4 early.	Project Manager			
9/22/04	Call placed informing 7-Day Safety	Ms. Tanima Sinha	N/A	N/A	2004-BP-08103BP(0)
	Report would be faxed.	Project Manager			
9/21/04	FDA Request for Desk copies –	M. Tanima Sinha	9/7/04	Original	Treatment IND
	Treatment IND	Project Manager		IND 70.629	
9/17/04	Response to Requests – August 30 <sup>th</sup>	Ms. Tanima Sinha	318	2/20/04	GC - Request for FDA Feedback in
	Telecon	Project Manager			regards to BI's plans for immunotoxicology testing for TPV
9/17/04	FDA Attendees for August 30th	Ms. Tanima Sinha	N/A	N/A	N/A
	telecon	Project Manager			

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9/13,16,20, 2004	FDA's Patient Profiler Text Files	Ken Edmunds FDA/CDER/OIM	N/A	N/A	N/A
9/3/04	TPV Electronic NDA Demonstration scheduled Oct. 5, 2004 10-11:30am	Ms. Tanima Sinha Project Manager	9/1/04	N/A	9/1/04 Telecon
	Submission of reports to the NDA  Treatment IND – Annual Report  Required				
9/3/04	BI Attendees for Sept. 1st Telecon	Ms. Tanima Sinha Project Manager	9/1/04	N/A	9/1/04 Telecon
9/3/04	FDA attendees from September 1st relecon	Ms. Tanima Sinha Project Manager	9/1/04	N/A	9/1/04 Telecon
9/3/04	Call placed informing 7-Day Safety Report would be faxed.	Ms. Tanima Sinha Project Manager	N/A	N/A	2004-BP-03153BR(0)
9/2/04	FDA contact regarding the 2-year rat carcinogenicity study 02R051	Dr. James Farrelly, Supervisory Pharmacologist, DAVDP	6/17/04	470	Information Amendment: PharmTox 24-Month Oral Rat Carcinogenicity Study
9/2/04	September 1st Telecon Clarification on in vitro interaction Data of TPV with APV and LPV	Ms. Tanima Sinha Project Manager	527	7/22/04	Information Amendment: Pharmacology/Toxicology (U03-3565, U03-3153)
9/1/04	Request for Information on NDA Logistics	Ms. Tanima Sinha Project Manager	N/A	N/A	N/A

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		PERSON		Reference	
8/30/04	Copies of Meeting Minutes or Faxes	Ken Edmunds	N/A	N/A	N/A
	Irom important FDA/Sponsor Meetings can be included in the	FDA/CDEK/OIM			
	literature sections of Modules 3 4				
	and 5.				
8/31/04	Call placed informing 7-Day Safety	Ms. Tanima Sinha	N/A	N/A	2004-BP-07145BP(0)
	Report would be faxed.	Project Manager			· · · · · · · · · · · · · · · · · · ·
8/27/04	Confirmed FDA/BIPI Teleconference	Mr. Destry	N/A	N/A	N/A
	time extended – 12:30-2:00 pm on 30	Sullivan			
	August 2004	Project Manager			
		FDA, CDER,			
		Division of			
		Antiviral Drug			
		Products			
8/26 & 8/31	FDA ESUB Coordinator provided	Ken Edmunds	N/A	N/A	N/A
	guidance on three questions related to	FDA/CDER/OIM			
	DVD's and their use as the media for		-		
	an electronic submission				
8/18/04	FDA Request for Time Change –	Destry Sillivan	N/A	N/A	N/A
	August 30th teleconference/resistance	Project Manager,			
	template/pharm tox	DAVDP			
8/16/04	Schedule Teleconference to Discuss	Destry Sillivan	N/A	N/A	N/A
	HIV Resistance Template	Project Manager,			
		DAVDF			
8/13/04	Location of Financial Disclosure	Ms. Tanima Sinha	N/A	N/A	N/A
	Information in NDA's	Project Manager			

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N/A	N/A	N/A
A/A	N/A	N/A
N/A	N/A	N/A
Gary Gensinger Supervisory Regulatory Info	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager
- Any electronic files that are not compliant with the file formats listed in the FDA's electronic submission guidance and submission guidance and submission before it is loaded onto FDA's Electronic Server  For the Tipranavir electronic submission Server For the Tipranavir electronic submission demo, BIPI should plan on bringing a laptop loaded with the data  - The esub coordination should be contacted with the request to "test load" electronic data	Follow-up Electronic Submission of Clinical Report 1182.4  • Request for Response on Outstanding Business	Electronic Submission of Clinical Report 1182.4 – BIPI requesting that appendices will be submitted electronically.
8/13/04	8/10/04	8/3/04

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		CONTACT PERSON	Reference	of Reference	
			•		
8/3/04	Delay in submission 1182.51 study	Ms. Tanima Sinha	4/15/04	383	Information Amendment –
	report to September 2004	Project Manager			Clinical
	Request for comment in preparation				Interim Study Results of
	for EAP telecon on August 4, 2004				Pharmacokinetic Study
	Follow up on outstanding business				1182.51
7/30/04	Updated HIV resistance template	Ms. Tanima Sinha	7/28/04	533	General Correspondence –
	Request for status update of feedback	Project Manager			Request for FDA Feedback
	on microbiology submissions				Resistance Template
	o Resistance template				
	o In vitro study proposal				
	Follow up on outstanding issues				
7/29/04	Call placed informing 7-Day Safety	Ms. Tanima Sinha	N/A	N/A	2004-BP-05975BP(0)
	Report would be faxed.	Project Manager			
7/28/04	DVD can be used as a media for	Ken Edmunds	N/A	N/A	N/A
	electronic submissions.	FDA/CDER/OIM			
7/27/04	Obtain clarification of User Fee for	Ms. Beverly	N/A	N/A	N/A
	upcoming NDA's for the same drug	Friedman			
	but different dosage form to be	Office of			
	sumitted at the same time.	Regulatory Policy,			
		HFD-5			
7/27/04	Call placed informing 7-Day Safety	Ms. Tanima Sinha	N/A	N/A	2004-BP-05812BP(0)
	Report would be faxed.	Project Manager			
7/22/04	Submission of New Report to NDA	Ms. Tanima Sinha	N/A	N/A	N/A
		Project Manager			
7/22/04	BI Follow up on IND Serial	Ms. Tanima Sinha	483	6/22/04	Type A IND Meeting-Response to
	Numbering	Project Manager			FDA Pre-Meeting Request for

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	Information-CMC-Tipranavir Oral Solution	Response to FDA Request for Information	Response to Telefax from FDA Pharmacologist, Anthony ElHage-	Requesting info on investigators, sites, discontinuations and reasons	discontinuations for all of the	pivotal sties for 1FV 1ffals   (1182.12, 1182.48 and 1182.14	General Correspondence /Request	for FDA Feedback /Request for	Teleconference	NDA Clinical Summary Cross-	Referencing Plan	Presentation of Patient	Disposition, Clinical Summary y	Integration Plan	Information Amendment - Clinical	Safety Information	• 2004-FF-00088FF(0)
Reference		513	6/3/04				501								315		
		7/12/04	telefax				7/1/04								2/18/04		
PERSON		Ms. Tanima Sinha Project Manager					Ms. Tanima Sinha	Project Manager								Project Manager	
		<ul> <li>Correction of Request for Desk Copies</li> </ul>	<ul> <li>Confirmation of Teleconference on 3-step approach</li> </ul>				Teleconference Between BIPI	and FDA on July 19, 2004	Attendees						SN 315 (Clinical Information	Amendment-Satety) Desk Copies	
		7/20/04					7/16/04								7/16/04		

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	,						
		General Correspondence/Request	for FDA Feedback/Request for	Teleconference NDA Clinical	Sumary Cross Referencing Plan,	Presentation of Patient Disposition,	Clinical Summary Integration Plan
Reference		501					
		7/1/04					
PERSON		for Desk Copies - Ms. Tanima Sinha	Project Manager				
	•.	ednest	SN 501				
		7/13/04		-			

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		I LINDOIN		Neielelle	
7/12/04	Telcon to discuss E-submission and adequacy of CDISC 3.0	Ms. Tanima Sinha Project Manager,	3/17/04	346	Pre-NDA background document
	•	DAVDP, IT	6/18/04	471	E-submission plans update
			6/11/04	ACR	Request for feedback on e-sub plans
7/9/04	Submission of New Reports to NDA	Ms. Tanima Sinha Project Manager	Pre-Nda Meeting	6/2/04	E-mail follow up regarding submission of new reports to NDA
		,	Telecon	6/28/04	Discussion of New Reports being submitted to NDA.
7/9/04	Follow up on Submission SN	Ms. Tanima Sinha	7/1/04	501	General Corespondence/Request
	501-Scheduling of Telecon	Project Manager			for FDA Feedback /Request for
					I eleconference
					NDA Clinical Summary Cross-
					Referencing Plan
					Presentation of Patient
					Disposition, Clinical Summary
					integration rian
1/9/04	BI Attendees for July 12, 2004	Ms. Tanima Sinha	N/A	N/A	Teleconference to discuss
	teleconference	Project Manager			electronic submission proposal and
					request for information on the TPV NDA
7/9/04	In vitro interaction studies	Ms. Tanima Sinha	6/29/04	496	Information Amendment: Clinical -
	w/ARV's request for desk copies	Project Manager			Step 1 of Drug Interaction Study

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7/9/04	Proposal for in vitro studies with	Ms. Tanima Sinha	6/29/04	496	Information Amendment
	ARV's	Project Manager			Clinical – Step 1 of Drug
					Interaction Study
	Electronic Submission Proposal				
	Teleconference attendees		6/18/04	SN 346	Electronic Submission Proposal
7/8/04	BI Request for FDA Attendees for	Ms. Tanima Sinha	N/A	N/A	BI Request for FDA Attendess for
	Telecon	Project Manager			July 12, 2004 Telecon re: TPV Electronic Submission
40/L//	Request for Desk Copies of SN496	Ms. Tanima Sinha	6/30/04	496	General Correspondence Minutes
		Project Manager			of June 7, 2004 Teleconference
7/6/04	BI Request for Teleconference to	Ms. Tanima Sinha	6/30/04	496	General Correspondence Minutes
	discuss Step 1 of Drug Interaction	Project Manager			of June 7, 2004 Teleconference
	Analysis				
7/1/04	TPV IND Serial Numbering	Ms. Tanima Sinha	6/22/04	483	Type A IND Meeting Response.
	BIPI used SN 483 twice: FDA stated	Project Manager			
	not to change				
7/1/04	FDA Comments on SN 470	Ms. Tanima Sinha	6/17/04	470	Information Amendment:
	Termination of Dosing Group in Rat	Project Manager			PharmTox 24-Month Oral Rat
	Care Study				Carcinogenicity Study
	Scheduling Telecon for Step 1 of 3-				
	step DI Analysis				
6/30/04	Postponement of Tradename	Ms. Tanima Sinha	2/6/04	307	General Correspondence - Request
	Evaluation	Project Manager			for Evaluation of Trade name
6/29/04	Template for Resistance Data	Ms. Tanima Sinha Project Manager	N/A	N/A	Ms. Sinha sent latest template for
		Talloce transment			שוח שומופופטנו ז זוו פוווווווווו פוווווווווווווו

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6/28/04	Confirmation of Teleconference to	Ms. Tanima Sinha	N/A	N/A	Teleconference confirmation
	discuss Electronic Submission	Project Manager			
6/24/04-6/28/04	Follow up on outstanding business;	Ms. Tanima Sinha	6/17/04	470	Information Amendment:
	Procedural (IND SNs, NDA)	Project Manager			PharmTox 24-Month Oral Rat
	Number, Electronic				Carcinogenicity Study
	Submission Plan				
	Clinical (Patient Disposition				
	Plans for NDA, DI Analysis				
	3-step Approach, Integration				
	Plans, IND Annual Report/2-				
	month Safety Update Plan,				
	Clinical Summaries Cross-				
	Referencing Plan)				
	<ul> <li>Nonclinical (Rat Carc.,</li> </ul>				
	Immunotox, In vitro				
	Resistance, Resistance				
	Template/Presentation of				
	Microbiology in CTD)				

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Export Waiver	2004-BP-04608BP(0)	2004-FF-00364FF(0)	2004-UK-00526UK(0) RESENT	2004-BP-04500RA(0)	Information Amendment – Type A IND Meeting Package	May 10 <sup>th</sup> 2004 pre-NDA Telecon
	N/A	N/A	N/A	N/A	460	N/A
	N/A	N/A	N/A	N/A	6/11/04	N/A
Rosemary Johann- Liang, MD Medical Team Leader with Tanima Sinha, Project Manager and Andrea James, MD, Medical Reviewer	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Ms. Tanima Sinha Project Manager	Rosemary Johann- Liang, MD Medical Team Leader	General Division Voicemail
FDA Comment on Export Authorization Request/Protocol	Call placed informing 7-Day Safety Report would be faxed.	Call placed informing 7-Day Safety Report would be faxed.	Call placed informing 7-Day Safety Report would be faxed. Resend 6/10 fax	Call placed informing 7-Day Safety Report would be faxed.	Electronic Submission Specifics	FDA request for feedback on FDA request for additions to Electronic Submission Plan
6/18/04	6/17/04	6/17/04	6/17/04	6/16/04	6/14/04	6/11/04

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6/10/04	Call placed informing 7-Day Safety Report would be faxed.	Ms. Tanima Sinha Project Manager	N/A	N/A	2004-UK-00526UK(0)
6/10/04	FDA Project Manager Away from Office	Ms. Tanima Sinha Project Manager, DAVDP	N/A	N/A	N/A
6/3/04	FDA request for discussion of like analyses core cases 2004BP03721 and 2004BP03614BR at Monday, June 7, 2004 Teleconference	Ms. Tanima Sinha Project Manager, DAVDP	5/24/04	SN 431 & SN 432	Like analyses core cases 2004BP03721 and 2004BP03614BR
5/10/04, 6/2/04	Clinical /Nonclinical Pre-NDA Meeting (Telecon 5/10, Face to Face 6/2)	DAVDP	3/17/04	346	Pre-NDA Package
5/27/04	FDA request summary info regarding food effects on bioavailability of SEDDS formulation in humans	Ms. Tanima Sinha Project Manager, DAVDP	5/28/04	445	Information Amendment Clinical Food Effect
5/27/04	FDA reminder request of May 7 to summarize at the pre-NDA meeting the RESIST protocol and amendments and the impact of each amendment on the statistical analysis plan	Ms. Tanima Sinha Project Manager, DAVDP	3/17/04	346	General Correspondence – Pre- NDA Meeting Package Request for Teleconference to Discuss key clin ical, statistical and format topics prior to face-to-face meeting
5/27/04	FDA request- from Dr. Zhang, Clinical Pharmacology reviewer for Tipranavir regarding SEDDS formulation in humans	Ms. Tanima Sinha Project Manager, DAVDP	N/A	N/A	N/A

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5/27/04	SN 375 & 401 FDA requesting updated lab info, specifically bilimbing	Ms. Tanima Sinha Project Manager,	4/12/04	375	IND Safety Report Initial Report 2003-BP-03027BP(0)
		TO A DO	5/3/04	401	Follow-up 2003-BP-09788BR(3)
5/27/04	FDA Request-summarize RESIST protocol & amendments-impact of each amendment on the statistical	Ms. Tanima Sinha Project Manager,	3/17/04	346	General Correspondence – Pre- NDA Meeting Package
	analysis plan for TPV				Discuss key clinical, statistical and format topics prior to face-to-face meeting
5/21/04	Call placed informing 7-Day Safety Report would be faxed.	Ms. Tanima Sinha Project Manager, DAVDP	N/A	N/A	2004-BP-03787BP(0)
5/21/04	Safety Reporting Process teleconference scheduling on June 7, 2004 – Pre NDA meeting plans/submission of RESIST Data	Ms. Tanima Sinha Project Manager, DAVDP	5/21/04	427	General Correspondence – Request for Teleconference – Update of Safety Reporting Procedures
5/11/04	Follow up on submission SN 318/Immunotox Proposal	Ms. Tanima Sinha Project Manager, DAVDP	2/20/04	318	General Correspondence – Request for FDA Feedback in regards to BI's plans for immunotoxicology testing for Tipranavir
5/10/04	FDA out of office notification	Ms. Tanima Sinha Project Manager, DAVDP	N/A	N/A	N/A

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5/7/04	BI Attendees for May 7, 2004 teleconference	Ms. Tanima Sinha Project Manager, DAVDP	5/7/04	N/A	teleconference
5/7/04	FDA request to briefly summarize the original RESIST protocol amendment	Ms. Tanima Sinha Project Manager, DAVDP	3/17/04	346	General Correspondence – Pre- NDA Meeting Package Request for Teleconference to Discuss key clinical, statistical and format topics prior to face-to-face meeting
5/6/04	FDA Attendees for May 7, 2004 teleconference	Ms. Tanima Sinha Project Manager, DAVDP	N/A	N/A	FDA attendees for May 7, 2004 teleconference
5/5/04	Background Information for Friday May 7, 2004 Teleconference	Ms. Tanima Sinha Project Manager, DAVDP	4/15/04	384	Response to Request for Information/Request for Teleconference
5/4/04	FDA Request for Desk copies SN303/SN392	Ms. Tanima Sinha Project Manager, DAVDP	2/6/04	303	IND Annual Report – Reporting period 10/1/02-9/30/03 Response to Request for Information: Tabular Listing of Deaths
4/29/04	Follow up on Submission SN318- February 20, 2004 Immunotox Proposal	Tanima Sinha Project Manager DAVDP	2/20/04	318	General Correspondence – Request for FDA Feedback in regards to BI's plans for immunotoxicology testing for Tipranavir

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4/26/04	TPV FDA Teleconference to address	Ms. Tanima Sinha	4/22/04	392	Response to Request for
	responses to April 8, 2004 Fax	Project Manager, DAVDP			Information: Tabular Listing of Deaths
4/16/04	Telecon arrangements discuss	Tanima Sinha	4/15/04	383	Information Amendment: Clinical
	responses to April 8/04 Fax	Project Manager,			Interim study results of
	(feedback on interim results from	DAVDP			pharmacokinetic study 1182.51
	study 1182.51)	5			
4/9/04	Respond to request for info (PK Shidy 1182 51 Methods/Procedures	Ms. Tanima Sinha	4/15/04	383	Information Amendment: Clinical
	for a Proposal to examine the	DAVDP			pharmacokinetic study 1182.51
-	possible effects of medication on				
	HIV-related death in TPV trials				
4/9/04	Advise Division of plans to respond		4/8/04	FDA Fax	Fax regarding pharmacokinetic
	to April 8, 2004 fax	Project Manager,			Study 1182.51
		DAVDP	4/15/04	383	IA Clinical - Interim Study
					Results of pharmacokinetic
					Study 1182.51
4/8/04	Fax regarding Pharmacokinetic Study	Rosemary Johann-	N/A	N/A	Telephone contact – FDA
	. 107011	Medical Team			informing of a fax being sent
		Medical Jeam			regarding the Pharmacokinetic
		Leader (with			Study 1182.51
		Tanıma Sinha,			
		Project Manager			
		and Andrea James,			
		MD, Medical			
		Reviewer			

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General Correspondence – Background/Current Safety Reporting Processes and Procedures and Proposed updates to Safety Reporting Processes and Procedures Request for Teleconference to assure that BI can implement the new Processes and Procedures	Timing for Pre-NDA teleconference	Information Package for Type B Pre-NDA Meeting - CMC	2004-BP-02413AU(0) 2003-BP-10433BP(0)	Scheduling of Pre-NDA Meeting	General Correspondence: Statistical Analysis Plan – submitting the summary of the RESIST Protocol
369	Agency Contacts	349	N/A	Agency Contact	14
4/6/04	3/12/04 & 3/26/04	3/19/04	N/A	3/12/04	3/17/04
Ms. Tanima Sinha Project Manager, DAVDP	Ms. Tanima Sinha Project Manager, DAVDP	Ms. Tanima Sinha Project Manager, DAVDP	Ms. Tanima Sinha Project Manager, DAVDP	Ms. Tanima Sinha Project Manager, DAVDP	Ms. Tanima Sinha Project Manager
Mtg to discuss update to safety reporting processes and procedures (sn 369)/timing of pre-NDA Mtg&Telecon	Timing for Pre-NDA teleconference	FDA Request for Desk Copies of CMC Pre-NDA Meeting Package SN 349	Informed that 2 7-day safety reports would be faxed.	Scheduling of Pre-NDA Meeting	Request for desk copy of RESIST Protocol.
4/7/04	4/2/04	4/2/04	3/31/04	3/26/04	3/17/04

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Follow-up on previous discussions about setting up a meeting.	Ms. Tanima Sinha Project Manager	N/A	N/A	N/A
Attention fax about 7-day safety	Ms. Tanima Sinha	N/A	N/A	2004-BP-00176-FF(0)
reports being sent.	Project Manager			2004-BP-00953BP(0)
				2004-FF-00180FF(0)
				2004-BP-01840BP(0)
Attention fax about 7-day safety	Ms. Tanima Sinha	N/A	N/A	2004-BP-01767BP(0)
reports being sent.	Project Manager			2004-BP-10562BP(0)
Mortality Analysis Submission,	Ms. Tanima Sinha	3/12/04	340	Response to FDA Request - Review
Request for Teleconference, and Pre-	<ul> <li>Project Manager</li> </ul>			of Mortality Rates
NDA Meeting Timing.				Request for Teleconference to
				discuss this information, answer any
				questions and receive feedback
Inform FDA decision to hold separate		2/17/04	314	Request for Type B Meeting - CMC
pre-NDA meeting for oral solution.	Project Manager			Pre-NDA Meeting
Confirm that CMC pre-NDA meeting	<b></b>			
will be 2 hrs long, ask about new				
Chemistry Reviewer				
Advisory of sending an electronic	Ms. Tanima Sinha		276	Information Amendment - Clinical
copy of a submission providing a	Project Manager			
Mortality Assessment of the				Request for FDA Comments on
tipranavir RESIST program.				Special Protocol Assessment for
				Protocol 1182.12 (RESIST 1)

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Description of Reference	,	
Serial No.	of	Reference
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REASON FOR CONTACT		
DATE		

	Call placed informing 7-Day Safety Report would be faxed. Request from FDA regarding mortality rates and publication discussed at Feb 19, 2004 telecon. Request for desk copies of SN 307, Proposed Trade name submission and feedback on timing for response to trade name evaluation submission. Call placed informing 7-Day Safety Report would be faxed. Call placed informing 7-Day Safety Report would be faxed. Email attachment with BI Attendees for 2/19/04 Teleconference. FDA Attendees and agenda for Feb 19, 2004.	Ms. Tanima Sinha Project Manager Ms. Tanima Sinha DAVDP Ms. Tanima Sinha DAVDP Ms. Tanima Sinha DAVDP	N/A 2/6/04 2/6/04 N/A N/A 1/12/04 1/12/04	N/A N/A N/A N/A 278	2003-BP-04672BP(1) 2003-FF-00625FF(0) February 19, 2004 teleconference regarding mortality rates and publication General Correspondence – Request for Evaluation of Trade name  204-SW-00048SW(0)  204-FF-00105FF February 19, 2004 teleconference General Correspondence – Response to Statistical Comments to Protocol 1182.12 (RESIST 2) Amendment 2 IND Safety Report
回ッズ	Email requesting all future export waiver requests be sent to David Kelly rather than Michele Limoli.	David Kelly, International Affairs	1/20/04 N/A	288 N/A	IND Safety Report Export waivers

REASON FOR CONTACT NAME OF Date of Serial No.  CONTACT Reference of PERSON Reference
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CMC pre-NDA meeting package had been submitted and to get an idea when meeting to be held.  Call placed informing 7-Day Safety Report would be faxed.  FDA response to request for pre-NDA meeting for tipranavir.  Call placed informing 7-Day Safety Project Manager  Call placed information at DAVDP.  Report would be faxed.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding Acting Project Pre-NDA Meeting request and contact information in Nancy  McKay's absence.  Degrees/titles for BIPI participants of Ms. Virginia Behr 1/16/04 N/A Sold-104 Information from BI regarding the Ms. Virginia Behr 1/16/04 N/A Sold-104 Information from BI regarding the Ms. Virginia Behr 1/16/04 N/A Sold-104-104-104-104-104-104-104-104-104-104	2/18/04	To alert CSO that a request for a	Tanima Sinha.	2/17/04	314	Reguest for Type B Meeting – CMC
when meeting to be held.  Call placed informing 7-Day Safety Report would be faxed.  FDA response to request for pre- NDA meeting for tipranavir.  Call placed informing 7-Day Safety NDA meeting for tipranavir.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding Acting Project  BI's plans for monthly safety update report, request for feedback regarding Acting Project  Manager  Contact information in Nancy  Manager  Degreess/titles for BIPI participants of Ms. Virginia Behr Information from BI regarding the Acting Project  Manager  Manager  Si04/04  Information from BI regarding the Acting Project Manager  Acting Project Manager  Manager  Manager  Acting Project Manager  Manager  Acting Project Manager  Acting Project Manager  Manager  Manager  Acting Project Manager  Acting Project Manager  Manager  Acting Project Manager  Acti		CMC pre-NDA meeting package had	CSO FDA			Pre-NDA Meeting
When meeting to be held.  Call placed informing 7-Day Safety Report would be faxed.  The first meeting for tipranavir.  Call placed information at DAVDP.  Report would be faxed.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding for contact information in Nancy  Manager  Contact information in Nancy  Manager  Contact information in Nancy  Manager  Contact information in Nancy  McKay's absence.  Si04/04  Information from BI regarding the feleconference for feb 19, 2004.  Acting Project  Manager  Manager  Acting Project  Manager  Acting Project  Manager  Manager  Acting Project  Manager  Manager  Acting Project  Manager  Manager  Manager  Acting Project  Manager  Manager  Manager  Acting Project  Manager  Manager  Acting Project  Manager  Manager  Acting Project  Manager  Mana		been submitted and to get an idea				
Report would be faxed.  Project Manger  H. Call placed informing 7-Day Safety Report would be faxed.  H. FDA response to request for pre- NDA meeting for tipranavir.  Call placed informing 7-Day Safety Report would be faxed.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding Router information in Nancy  MCKay's absence.  Degrees/titles for BIPI participants of Ms. Virginia Behr McKay's absence.  Degrees/titles for BIPI participants of Ms. Virginia Behr January 16, 2004 meeting.  Manager  Si04/04  Information from BI regarding the deting Project Router Peb 19, 2004.  Manager  Acting Project Manager  Acting Project Manager  Acting Project Manager  Acting Project Manager		when meeting to be held.				
Report would be faxed.  FDA response to request for pre- NDA meeting for tipranavir.  Call placed information at DAVDP.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding Project Pre-NDA Meeting request and contact information in Nancy  Manager  Degrees/titles for BIPI participants of Ms. Virginia Behr JJ30/04  Degrees/titles for BIPI participants of Ms. Virginia Behr JJ30/04  Degrees/titles for BIPI participants of Ms. Virginia Behr JJ30/04  January 16, 2004 meeting.  Manager  S/04/04  Information from BI regarding the Acting Project Robins Source Acting Project Manager	2/13/04	Call placed informing 7-Day Safety	Tanima Sinha,	N/A	N/A	2004-BP-01039BP(0)
FDA response to request for pre- NDA meeting for tipranavir.  Call placed informing 7-Day Safety Report would be faxed.  Contact information at DAVDP. BI's plans for monthly safety update report, request for feedback regarding Project Pre-NDA Meeting request and contact information in Nancy  MCKay's absence.  Degrees/titles for BIPI participants of MS. Virginia Behr 1/16/04 January 16, 2004 meeting.  Degrees/titles for BIPI participants of MS. Virginia Behr 1/16/04 Manager  Manager  Acting Project Manager  Acting Project Manager  Manager  Acting Project Manager		Report would be faxed.	Project Manger			2004-FF-00088FF(0)
FDA response to request for pre- NDA meeting for tipranavir. Call placed informing 7-Day Safety Report would be faxed. Contact information at DAVDP. BI's plans for monthly safety update report, request for feedback regarding Project Manager contact information in Nancy Manager Contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of Ms. Virginia Behr 1/16/04 January 16, 2004 meeting. Ms. Virginia Behr 1/16/04 Manager S/04/04 Information from BI regarding the Acting Project Feb 19, 2004.	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,					2004-BP-00954BP(0)
NDA meeting for tipranavir.  Call placed informing 7-Day Safety Report would be faxed. Contact information at DAVDP. Bl's plans for monthly safety update report, request for feedback regarding Rochact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Degrees/titles for BIPI participants of Ms. Virginia Behr January 16, 2004 meeting. Manager	2/12/04	FDA response to request for pre-	Ms. Tanima Sinha	1/30/04	297	Request for Type B Meeting - Pre-
Call placed information at DAVDP.  Report would be faxed.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding Project Pre-NDA Meeting request and contact information in Nancy  Manager  Manager  Acting Project Manager  Manager  Contact information in Nancy  Manager  Degrees/titles for BIPI participants of Ms. Virginia Behr Jamuary 16, 2004 meeting.  Manager  Manager  Manager  Jamuary 16, 2004 meeting.  Ms. Virginia Behr Ms. Virginia Behr Ms. Virginia Behr Jamuary 16, 2004 meeting.  Manager  Acting Project  Manager  Acting Project  Manager  Acting Project  Manager  Feb 19, 2004.		NDA meeting for tipranavir.	Project Manager			NDA meeting request
Report would be faxed.  Contact information at DAVDP.  Contact information at DAVDP.  BI's plans for monthly safety update report, request for feedback regarding Project Pre-NDA Meeting request and contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of Ms. Virginia Behr 1/16/04 N/A January 16, 2004 meeting.  Manager Manager Acting Project Manager Manager Acting Project Manager Manager Acting Project Manager Feb 19, 2004.  Manager Feb 19, 2004.	2/9/04	Call placed informing 7-Day Safety	Tanima Sinha,	N/A	N/A	2004-BP-00825BR
Contact information at DAVDP. Ms. Virginia Behr N/A Acting Project Manager  BI's plans for monthly safety update Rs. Virginia Behr report, request for feedback regarding Acting Project Pre-NDA Meeting request and contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of Ms. Virginia Behr I/16/04 N/A January 16, 2004 meeting. Manager Manager Manager Feb 19, 2004.  Feb 19, 2004.  Manager Acting Project Acting Project Feb 19, 2004.		Report would be faxed.	Project Manager			2004-IT-00012IT
BI's plans for monthly safety update report, request for feedback regarding Acting Project Pre-NDA Meeting request and contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Manager  S/04/04 Information from BI regarding the teleconference for Feb 19, 2004.  Manager  Acting Project Manager  I/16/04 N/A Acting Project Manager  Acting Project Manager  Acting Project Manager  Manager  Feb 19, 2004.	2/6/04	Contact information at DAVDP.	Ms. Virginia Behr	N/A	N/A	N/A
Manager BI's plans for monthly safety update report, request for feedback regarding report, request for feedback regarding Acting Project Pre-NDA Meeting request and Manager contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Manager  Manager  Manager  Manager  1/16/04  N/A  Acting Project Manager  Manager  Acting Project Feb 19, 2004.  Manager  Manager			Acting Project			
Bl's plans for monthly safety update report, request for feedback regarding Acting Project Pre-NDA Meeting request and contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Manager  5/04/04 Information from BI regarding the releconference for Feb 19, 2004.  Banager  Ms. Virginia Behr  1/7/04  299  1/30/04  297  1/30/04  297  Acting Project  Ms. Virginia Behr  Ms. Virginia Behr  Acting Project  Manager  Acting Project  Ranager  Manager			Manager			
report, request for feedback regarding Project Pre-NDA Meeting request and contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Manager  S/04/04 Information from BI regarding the teleconference for Feb 19, 2004.  Manager  Acting Project Manager  Manager  Manager	2/6/04	BI's plans for monthly safety update	Ms. Virginia Behr	2/2/04	299	General Correspondence - drafted
Pre-NDA Meeting request and Contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Manager  Mother Information from BI regarding the teleconference for Feb 19, 2004.  Manager  Manager  Manager  Manager  Manager		report, request for feedback regarding	Acting Project			revision to corporate Standard
contact information in Nancy McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  Manager  Information from BI regarding the teleconference for Feb 19, 2004.  Manager  Solution from BI regarding the Acting Project Acting Project  Manager		Pre-NDA Meeting request and	Manager			Operating Procedures (SOPs)
McKay's absence.  Degrees/titles for BIPI participants of January 16, 2004 meeting.  January 16, 2004 meeting.  Manager  Mormation from BI regarding the teleconference for Hanager  Feb 19, 2004.		contact information in Nancy				regarding adverse event reporting
Degrees/titles for BIPI participants of January 16, 2004 meeting.  January 16, 2004 meeting.  Manager  S/04/04  Information from BI regarding the teleconference for Feb 19, 2004.  Manager  Manager  N/A  N/A  N/A  Acting Project  Acting Project  Manager		McKay's absence.		1/30/04	297	Request for Type B Meeting - Pre-
Degrees/titles for BIPI participants of Ms. Virginia Behr 1/16/04 N/A January 16, 2004 meeting. Acting Project Manager  S/04/04 Information from BI regarding the teleconference for Hoting Project Feb 19, 2004. Manager						NDA meeting request
January 16, 2004 meeting.  Manager Information from BI regarding the teleconference for Acting Project Feb 19, 2004.	2/5/04		Ms. Virginia Behr	1/16/04	N/A	January 16, 2004 meeting
Information from BI regarding the Ms. Virginia Behr 1/7/04 278 teleconference for Acting Project Feb 19, 2004.		January 16, 2004 meeting.	Acting Project			
Information from BI regarding the Ms. Virginia Behr 1/7/04 278 teleconference for Acting Project Feb 19, 2004.			Manager			
Acting Project Manager	2/3/04-5/04/04	_	Ms. Virginia Behr	1/7/04	278	General Correspondence - Response
Manager		teleconference for	Acting Project			to Statistical Comments to Protocol
		Feb 19, 2004.	Manager			1182.12 (RESIST 1) Amendment 2

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OATE	REASON FOR CONTACT	NAME OF	Date of	Serial No.	Description of Reference	
		CONTACT	Reference	Jo.		
		PERSON		Reference		

2/2/04	<ul> <li>Call placed informing 7-Day Safety Report would be faxed.</li> <li>Also, Virginia gave new fax number.</li> </ul>	Ms. Virginia Behr Acting Project Manager	N/A	N/A	Case # 2003-BP-00677BP(0)
2/2/04	the safety mission. ference to ag and creport creport stical	Ms. Virginia Behr Acting Project Manager	2/2/04	299	General Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting
1/30/04	Status of BI's progress with agreements made at Jan 16, 2004 meeting.	Ms. Virgina Behr Acting Project Manager	2/5/04	302 telecon	General Correspondence – BI Meeting minutes from face to face meeting – January 16, 2004 Teleconference regarding Safety reporting
1/28/04	BIPIs electronic presentations from 1/16/04 face-to-face meeting. Request for copy of FDA presentation.	Virginia Behr, DAVDP	N/A	N/A	BIPIs electronic presentations from 1/16/04 face-to-face meeting. Request for copy of FDA presentation
1/27/04	<ul> <li>Call placed informing 7-Day</li> <li>Safety Report would be faxed.</li> </ul>	Virgin ia Behr, DAVDP	N/A	N/A	Case # 2003-BP-08944BP

of	Reference
Reference	
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	Reference

- 26 RTV 3 week			or ss and		- 26 RTV 3 week
Information Amendment – Pharmacology/Toxicology – 26 week Safety Study of TPV/RTV SEDDS in Beagle Dogs – 13 week draft interim report	Tox data	N/A	Response to FDA request for information – Safety Policies and TPV Risk Benefits	FDA telecon	Information Amendment – Pharmacology/Toxicology – 26 week Safety Study of TPV/RTV SEDDS in Beagle Dogs – 13 week
255 II	Agency T	N/A	285 F	N/A F	255 I
9/26/03	1/13/04	N/A	1/14/04	1/9/04	9/26/03
Ms. Virginia Behr Acting Project Manager		Virginia Behr, DAVDP	Virgina Behr, DAVDP	Virgina Behr, DAVDP	Ms. Virginia Behr Project Manager
FDA inquiry about status of Cremphor EL study and request for slides for meeting presentations.		Email attachment with FDA attendees for 1/16/04 face-to-face meeting.	FDA confirmation of receipt of SN 285.	FDA Attendees for 1/9/04 teleconference	Toxicology data for meeting between BI and FDA on January 16, 2004.
1/15-16/04		1/15/04	1/14/04	1/13/05	1/13/04

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Response to FDA request for information-providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxyl 135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials	General Correspondence Safety Information for January 8, 2004 Telecon	Response to Request for Information  - Aes and SAEs	Teleconference regarding Safety reports from Abbott on RTV	General Correspondence Safety Information for January 8, 2004 Telecon
150	277	279	N/A	277
5/17/02	1/7/04	1/9/04	1/8/04	1/7/04
Ms. Virginia Behr Project Manager	Virginia Behr, DAVDP	Virginia Behr, DAVDP	Virgina Behr, DAVDP	Virginia Behr, DAVDP
FDA called to inquire about the status Ms. Virginia Behr of the 26 week dog study.  Project Manager	Change of reporting period for the FDA request for Safety Information	Status of SN 279 Response to FDA Request for Safety Information.	Names, titles and degrees for FDA and BIPI participants of 1/8/04 teleconference.	FDA Request for information-AEs and SAEs (change in due dates of submissions).
1/12/04	1/9/04	1/9/04	1/8/04	1/8/04-1/9/04

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1/10/03	FDA called to ask where tinranavir	Ms. Bettv McRov	N/A	N/A	Export waivers
	capsules were manufactured for purposes of export waiver review.				
11/7/03	FDA Request for long-term results for 1182.52.	Tanima Sinha Project Manager	N/A	N/A	N/A
10/27/03-10/30/03	Email regarding export waiver clarification and request from microbiology team leader.	Tanima Sinha, Div. of Antiviral Drug Products	N/A	N/A	Export waivers
10/24/03	Confirmation that desk copies of the naïve special protocol assessment will be sent today.	Tanima Sinha, Project Manager	9/26/03	254	Request for Special Protocol Assessment – Clinical, Naïve Trial 1182.33
	Request clarification regarding hold on export waiver requests for 1182.14 study.				Export waivers
10/24/03	Request for summary on in vitro selection on virus resistance to TPV.	Tanima Sinha, Project Manager	11/20/03	265	Information Amendment – Clinical, Response to FDA Request for information On in vitro Selection of Virus Resistant to Tipranavir

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Export waiver – previous request for status update	Export waiver – previous request for status update			ndate	annd <sub>n</sub>										
aiver – pr late	aiver – pr late	aiver	aiver.	Reguest for Status undate	Contract of the Contract of th		'aiver								
Export waive status update	Export waive status update	Export waiver	Export waiver.	Regueet f	r andray.		Export Waiver	1							
Agency Contact	Agency Contact	N/A	N/A	Agency	Contact		N/A								
2/03	03-	03	03							•					
10/16/03	10/8/03-	8/20/03	8/20/03	10/9/03-	10/8/03	5	8/20/03								
Tanima Sinha, DAVDP			Tanima Sinha,	DAVDP			Tanima Sinha,	DAVDP, Betty	McRoy, OC, David	Kelly, IAS,	Elizabeth Duvall,	OND			
Email regarding status of review for Tipranavir Export Waiver Request for 1182 14. Argenting Brazil	Mexico, and Russia.		status	of review for Tipranavir Export Waiver Regulest for 1182-14-	Argentina, Brazil, Mexico, and	Russia.	r export		Argentina, Brazil Mexico and Russia		o o	Norris and Mr. David Kelly in IAS			
10/20/03			10/16/03				10/06/03-10/08/03								

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	Export waiver for 1182.17 Argentina	Protocol Amendment – New Investigators 1182.12-1182.17- 1182.24-1182.51 Administrative Info for Telecon Administrative Info for Telecon Administrative Info for Telecon Pediatric Written Agreement	Pediatric Written Agreement	Confirmation of teleconference and data – October 2, 2003, 1-2:00pm	Pediatric Written Agreement Admin. Info for telecon Confirm date for Telecon
COLOTOTO	N/A	250 ACR's (2) ACR ACR SN 242	242	Agency Contact	242 ACRs (2) ACR
	6/6/03	9/12/03 9/24/03 9/22/03 9/17/03 7/23/03	7/23/03	9/17/03 & 9/22/03	7/23/03 9/17/03 8/27/03
NOCKET I	Betty McRoy, OC Anna Noris, IAS	Tanima Sinha, Project Manager	Tanima Sinha, Project Manager	Tanima Sinha, Project Manager	Tanima Sinha, Project Manager
	Status of approval for tipranavir export waiver request for 1182.17-Argentina.	Missing pages from New Investigator Submission. Confirmation of teleconference info. for Pediatric written agreement telecon.	Administrative information for the telecon to discuss BI's Pediatric Written Agreement. Call in info & BI attendee list for Oct. 2, 2003 telecon	Confirmation to reschedule telecon for proposed Pediatric Written Agreement time of 10/2/03, 1-2 pm. Advised that Call-In info. Remained same for telecon.	Telecon to discuss BI Pediatric Written Agreement date/time negotiation.
	9/26/03-9/29/03	9/24/03	9/24/03	9/24/03	9/22/03

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	Pediatric Written Agreement Confirm date for Telecon	N/A	Export waiver – 1182.17 for Argentina	Request for status of export waiver				
Kelerence	242 ACR	N/A	N/A	ACR	ACR	ACR	ACR	ACR
	7/23/03 8/27/03	N/A	6/6/03	6/8/6-20/9/6	8/23/03- 8/29/03	8/6/03-8/8/03	8/12/03- 8/15/03	7/31/03-
FERSOIN	Tanima Sinha, Project Manager	Tanima Sinha, Project Manager	Tanima Sinha, DAVDP Betty McRoy, OC	Beth Duvall, OND				
	Call-In Information for the teleconference to discuss BI's Proposed Pediatric Written Agreement.	BI Attendees for teleconference on 9/18/03.	Status of Tipranavir Export Waiver Request for 1182.17-Argentina.					
	9/17/03	9/17/03	9/12/03-9/15/03					

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	waiver	waiver waiver	waiver waiver waiver	waiver waiver waiver	waiver waiver waiver	waiver waiver waiver	waiver waiver waiver	waiver waiver waiver	waiver waiver waiver	waiver waiver
Export waiver – 1182.17 for Argentina	Regilest for status of export	Request for status of export waiver Request for status of export waiver	Request for status of export waiver Request for status of export waiver Request for status of export waiver	Request for status of export waiver Request for status of export waiver Request for status of export waiver	Request for status of export vequest for export vequest for status of export vequest for e	Request for status of export v Pediatric Written Agreement	Request for status of export vequest for e	Request for status of export vequest for telecon	Request for status of export v Pediatric Written Agreement Proposed date of telecon Request for telecon Pediatric Written Agreement	Request for status of export vequest for status of export vequest for status of export vequest for status of export vediatric Written Agreement Proposed date of telecon Request for telecon Pediatric Written Agreement Request for telecon Request for telecon
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9/9/93	8/23/03-	8/23/03- 8/29/03 8/6/03-8/8/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03 7/31/03- 8/15/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03 7/31/03- 8/15/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03 7/31/03- 8/15/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03 7/23/03 8/26/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03 7/23/03 8/26/03 8/15/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/15/03- 8/15/03- 8/15/03 8/26/03 8/15/03	8/23/03- 8/29/03 8/6/03-8/8/03 8/12/03- 8/15/03 7/31/03- 8/15/03 8/26/03 8/15/03 8/15/03 8/15/03
Betty McRoy, OC					Betty McRoy, OC	Betty McRoy, OC Tanima Sinha, Project Manager	Betty McRoy, OC Tanima Sinha, Project Manager	Betty McRoy, OC Tanima Sinha, Project Manager	Betty McRoy, OC Tanima Sinha, Project Manager Tanima Sinha,	Betty McRoy, OC Tanima Sinha, Project Manager Tanima Sinha, Project Manager
Status of Tipranavir Export Waiver Request for 1182.17-Argentina.					Status of Tipranavir Export Waiver.	Status of Tipranavir Export Waiver. FDA confirms date for telecon to discuss BI Pediatric Written	Status of Tipranavir Export Waiver.  FDA confirms date for telecon to discuss BI Pediatric Written  Agreement – 9/18/03 11 am	Status of Tipranavir Export Waiver. FDA confirms date for telecon to discuss BI Pediatric Written Agreement – 9/18/03 11 am	Status of Tipranavir Export Waiver.  FDA confirms date for telecon to discuss BI Pediatric Written  Agreement – 9/18/03 11 am  Email with proposal dates for planned telecon between BI and FDA	Status of Tipranavir Export Waiver. FDA confirms date for telecon to discuss BI Pediatric Written Agreement – 9/18/03 11 am Email with proposal dates for planned telecon between BI and FDA regarding BI's pediatric written agreement.
9/6/03-9/8/03					9/5/03-8/31/03	9/5/03-8/31/03	9/5/03-8/31/03	9/5/03-8/31/03	9/5/03-8/31/03	9/5/03-8/31/03

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		Export Waiver 1182.17 Argentina	Request for status of export waiver	Request for status of export waiver	Request for status of export waiver	Pediatric Written Agreement	Resist 1 Protocol Amendments	Amendments 1&2	Amendment 3	Amendment 4	Pediatric Written Agreement		Export Waiver 1182.17 Argentina	Request for status export waiver	Request for status of export waiver
of Reference		N/A	ACR	ACR	ACR	242	214		220	230	242		N/A	ACR	ACR
Reference		20/9/9	8/6/03-8/8/03	8/12/03- 8/15/03	7/14,15,23, 29/03	7/23/03	3/19/03		3/31/03	6/10/03	7/23/03		6/6/03	8/6/03-8/8/03	7/14,15,23, 29/03
CONTACT		Betty McRoy, OC	CCI (SIIIONI PIIIIN			Tanima Sinha, Project Manager	Tanima Sinha,	Project Manager					Tanima Sinha DAVDP	Betty McRoy, OC	
		Status of approval for Tipranavir	Argentina			Request for telecon to discuss BI Pediatric Written Agreement.	Request of additional copies	of the RESIST 1 protocol	amendments	Inquired about telecon	between BI and FDA reg:	BI's written agreement for pediatric studies	Status of Tipranavir Export Waiver Request - 1182.17 Argentina	•	
		8/25/03-8/29/03				8/19/03-8/21/03	8/15/03						8/12/03-8/15/03		

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8/6/03-8/8/03	Status of Tipranavir Export Waiver	Betty McRoy	6/6/03	N/A	Export Waiver 1182.17 Argentina
	Nequest - 1102.17 Atgentula	Compliance Tanima Siha	7/31/03-8/5/03	ACR	Request for status of export waiver
		DAVDP	7/14,15,23, 29/03	ACR	Request for status of export waiver
7/31/03-8/5/03	Status of Tipranavir Export Waiver – 1182.17 Argentina.	Betty McRoy, Office of	6/6/03	N/A	Export Waiver 1182.17 Argentina
		Compliance	7/14,15,23, 29/03	ACR	Request for status of export waiver
7/18/03	Mechanism for submission of naïve protocol for FDA feedback	Tanima Sinha, Project Manager	11/15/02	186	General Correspondence – Background Document for End of
	Discussed whether the protocol should be submitted.				Phase II Meeting
	as a Special Protocol		7/15/03	ACR	Request for confirmation of Spec
	Assessment or an Information Amendment				Protocol Assess as mechanism for review of naive protocol.
	Response to pediatric written request		7/23/03	242	Proposed Changes in Written
	planned; teleconference to be requested				Request for Pediatric Studies/Proposed Written
	Submission to be sent week				Agreement for Pediatric
	of 21 July 03				Studies\Request for Teleconference
	BIPI requests a telecon be				
	scheduled to discuss the				
	written request				

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		LCCUT		Veleicilce	
7/15/03	Request for confirmation of Special	Tanima Sinha,			
	Protocol Assessment as Mechanism	Project Manager			
	for review of Naïve Protocol.				
7/14,15,23,29/03	Requesting Status of Export Waiver	Anna Norris, IAS	6/6/03	N/A	Export Waiver - 1182.17 Argentina
	Requests for 1182.17 - Argentina	Betty McRoy, OC			
6/10/03-6/24/03	Email re: information on patient	Jane Scott, PhD	N/A	N/A	E-mail correspondence
	reported outcomes.				
6/5/03	Request status of approval for	Anna Norris	3/12/03	N/A	Export Waiver - 1182.48
	1182.48 (Argentina)	Office of			Argentina
		International	5/8/03	ACR	DAVDP's review of export
		Affairs			waiver
			4/8/-11/03	ACR	Export Waiver Status
6/2-4/03	Request status of export waiver	Anna Norris	3/12/03	N/A	Export Waiver Request -
	requests for 1182.48 for Mexico	Office of			Brazil 1182.48
	(submitted on March 14, 2003) and	International	3/14/03	N/A	Export Waiver Request - Mexico
	Brazil (submitted on March 12,	Affairs			1182.48
	2003).		5/8/03	ACR	DAVP Review Status of export
				-	Waiver
			4/8-11/03	ACR	Export waiver status
5/21/03	Status of the CAC review of mouse	Tanima Sinha,	3/24/03	216	Request for Special Protocol
	carcinogenicity study	Project Manager			Assessment-Carcinogenicity Study
					Protocol. BI asking FDA
					concurrence with the mouse
					carcinogenicity study protocol
					= = = = = = = = = = = = = = = = = = = =
_	_		5/70/03	ACK	Request for Feedback on SN 216

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Request for Special Protocol Assessment-Carcinogenicity Study Protocol. BI asking FDA concurrence with the mouse carcinogenicity study protocol	SN Discrepancy	EoP2 Mtg Minutes (BIPI)	FDA EoP2 Mtg Minutes		1182.51 Protocol	Protocol Amendments 1&2 1182.51									
216	ACR	217	N/A	700	 402	526									
3/24/03	4/28/03	3/24/03	3/14/03	50,00	1/30/03	5/9/03									
Tanima Sinha, Project Manager	Virginia Yoerg, Regulatory Health	Project Manager	Tanima Sinha,	Regulatory Health	Project Manager										
Request for feedback on SN 216 Request for SPA – Carcinogenicity Protocol submitted 3/24/03.	DAVDP asked that BI resolves     serial number discrepancy directly	with document control room	<ul> <li>Introduction of new project manager, Tanima Sinha</li> </ul>	<ul> <li>Full or draft protocols should be</li> </ul>	included in export waiver request when possible	<ul> <li>DAVDP's review of export waiver</li> </ul>	requests for 1182.48	(Argentina/Mexico) is complete and	has been further processed within	FDA	<ul> <li>Follow-up responses on FDA</li> </ul>	comments on 1182.51 protocol	<ul> <li>Medical reviewer denied the need</li> </ul>	for a change in the official EOP II	Meeting Minutes
5/20/03	5/8/03														

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5/7/03	DAVDP clarified that it is not	Virginia Yoerg,	N/A	N/A	Export waiver
	necessarily mandatory to submit the	Regulatory Health			•
	protocols in their entirety when	Project Manager			
	request export waivers, however, it is				
	strongly encouraged for all pivotal				
	trials.				
5/5/03	DAVDP requested full copy of	Virginia Yoerg,	11/15/02	186	General Correspondence –
	RESIST 2 protocol in order to review	Regulatory Health			Background Document for end of
	BI's request for export waivers for	Project Manager			Phase II Meeting
	1182.48				,
	Serial Numbers discrepancy is still		4/22-4/28	ACR	DISCREPENCY WITH SERIAL
	under review.		4/18/03	ACR	NO.'S
4/28/03	Email to FDA clarifying serial	Virginia Yoerg,	4/22-4/28	ACR	DISCREPENCY WITH SERIAL
	number discrepancy.	Regulatory Health	4/18/03	ACR	NO.'S
		Project Manager			
4/28/03	FDA faxed over serial number 212	Virginia Yoerg,	4/22/-4/28	ACR	DISCREPENCY WITH SERIAL
	for clarification of serial number	Regulatory Health	4/18/03	ACR	NO.'S
	discrepancy.	Project Manager			
	EMail requesting agreement in that				
	letter should have not received a				
	serial number.	-			
4/22/03	Export Authorization Request Status	Andrea Chen,	N/A	N/A	N/A
	for IAS-D-3-3-17 and IAS-D-3-3-18.	Office of New			
		Drugs			

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4/22-28/03	Clarification of Serial Number	Virginia Yoerg,	4/18/03	ACR	DISCREPENCY WITH SERIAL
	discrepancy.	Regulatory Health Project Manager			NO.'S NO SUBMISSION
4/18/03	Serial Number discrepancy: FDA's serial numbers differ from BI's.	Virginia Yoerg, Regulatory Health	N/A	N/A	DISCREPENCY WITH SERIAL NO.'S
	Export Waiver Request Review	Project Manager			NO SUBMISSION
	Status: Medical reviewer currently reviewing export authorization				
	request.				
4/14/03	Export Waiver Status/Review of	Betty McRoy,	4/14/03	N/A	TPV Export Waiver Status/Review
	FDA Internal. Export Waiver Process	Office of	4/8/03	ACR	of FDA internal export waiver
	and Timing.	Compliance	4/11/03	ACR	review process and timing. #IAS-D-
		Andrea Chen,			3-3-17/IAS-D-3-3-18
		Office of New			
		Drugs			
4/8-11/03	Export Waiver Approval for 1182.48.	Michelle Limoli			
	(Mexico/Argentina) Status –	and Anne Norris			
	Expected in late May/early June.	FDA, International			
		Affairs			
3/27/03	Pam C. followed up on the 12/11/02	Virginia Yoerg,	12/11/02	191	General Correspondence-chemistry,
	submission reg: designation of	Regulatory Health			manufacturing and controls, BI
	starting materials and on action item	Project Manager			providing current specifications for
	form the EOPII meeting reg:				2 TPV and information regarding
	designation of Ph. Eur. Procedures as				M-nitropropriophenone (m-NPP)
	alternate procedures.				

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		PERSON		Reference	
3/21/03	Left voice mail with FDA re design	Virginia Yoerg,	2/7/03	206	General Correspondence - Request
	of the drug interaction study with	Regulatory Health			for FDA Feedback regarding drug
	atorvastatin (SN 206). Call was	Project Manager			interaction study between Tipranavir
	returned advising draft reviews				and Atorvastin
	almost completed & that they would				
	contact us when finished.		3/5/03	ACR	Comment on Atorvastatin Study
3/5/03	Called FDA for 1) clarification of	Virginia Yoerg,	2/7/03	206	General Correspondence - Request
	SPA of carcinogenicity protocols	Regulatory Health	2/6/03	ACR	for FDA Feedback regarding drug
	within the DAVDP and 2) for	Project Manager			interaction study between Tipranavir
	comment on S/N 206 re design of the	1			and Atorvastin
	drug interaction study with	, , .			
	atorvastatin.			•	
3/4/03	Verbal response from FDA approving	Virginia Yoerg,	12/13/02	196	General Correspondence - Request
	our proposal to study oral	Regulatory Health			for FDA Feedback regarding
	Cremophor.	Project Manager			proposal for the study of oral
					Cremophor EL human exposure
			9		
			2/6/03	ACR	Status request on feedback for CrEl
					human exposure
2/26/03	Email attachment sent to request	Virginia Yoerg	1/24/03	203	General Correspondence
	reedback on proposed changes.	Regulatory Health			
	Phone call was also made.	Project Manage r			

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2/6/03	Status of S/N 196. Waiting for	Virginia Yoerg,	12/13/02	196	General Correspondence - Request
	written response from FDA to our	Regulatory Health			for FDA Feedback regarding
	proposal to study oral Cremophor EL	Project Manager			proposal for the study of oral
	human exposure.				Cremophor EL human exposure
1/16/03	Pharm/Tox Teleconference - FDA	Virginia Yoerg,	1/15/03 &	Agency	Scheduling of Pharm/Tox
	will advise need for call next week.	Regulatory Health	1/3/03	Contact	teleconference
		Project Manager			
1/15/03	Pharm/Tox Teleconference - Still	Virginia Yoerg,	1/3/03	Agency	Scheduling of telecon
	attempting to schedule telecon to	Regulatory Health		Contact	
	discuss the Toxicology study of	Project Manager			
	degradants				
1/6/03	List of Attendees for EOP2 Clinical	Nitin Patel,	12/17/02	N/A	EOP2 Clinical meeting
	Meeting.	DAVDP, FDA			)
1/3/03	Pharm/Tox Teleconference requested	Virginia Yoerg,	N/A	N/A	N/A
	by FDA – Offered BI's availability	Regulatory Health			
	for dates/times on either Jan. 6, 7 and	Project Manager			
	∞				
1/3/03	1/3/03 fax re: Microbiology	Virginia Behr,	11/15/02	186	Phenotypic Assay
	comments SN 186-Phenotypic Assay.	DAVDP			
12/30/02	Protocol for the Animal Safety Study	Dr. Nitin Patel,	12/13/02	Telecon	Protocol for Animal Safety in dogs
	in Dogs – It BIPI has urgency tor	Project Manager			
	review, contact Ms. Yoerg and send				
	protocol via e-mail. Mr. Patel is out				
	of office until Jan. 6, 2003.				• 1

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12/26/02	Pharm/Tox Teleconference requested	Dr. Nitin Patel,	N/A	N/A	N/A
	during week of Jan. 6, 2003 to discuss the study of degradants.	Project Manager			
12/18/02	CMC EOP 2 Meeting.	N/A	12/13/02	194	General Correspondence –
					Background Document for End of Phase 2 meeting
12/17/02	Clinical/Nonclinical End of Phase 2	Dr. Nitin Patel	11/15/02	186	End of Phase 2 Meeting
	meeting	Project Manager			Package
12/13/02	FDA Contact Information for Dr.	Dr. Nitin Patel,	N/A	N/A	N/A
	Gitternan.	Project Manager			
12/12/02	Pharm/Tox Comments on 26 week	Dr. Nitin Patel,			FDA fax –
	Safety Tox Study and	Project Manager			Pharmacology/Toxicology
	Teleconference.				comments in preparation for the
					12/13/02 teleconference
12/11/02	RESIST II Protocol. Cancel telecon scheduled for today (12/11/02)	Dr. Nitin Patel, Project Manager	N/A	N/A	N/A
12/10/02	Ask about structure of degradation	Dr. Boring,	12/13/02	194	General Correspondence –
	products specified in list of tests for	Chemistry			Background Document for End of
	drug product given in the EOP 2	Reviewer			Phase 2 meeting
,	package.				
12/10/02	Fax re: clinical and clinical	Dr. Nitin Patel,	11/15/02	186	General Correspondence-
	prominaciogy commons (EQ1.2)	i ojeci ivianagei			Phase II Meeting

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12/9/02	trative	Dr. Nitin Patel,	12/9/02	Agency	Telecon schedule
	details for Pharm/Tox Telecon	Project Manager		Contact	
	scheduled for 12/11/02.				
12/9/02	EMail correspondence discussing	Dr. Nitin Patel,	N/A	N/A	N/A
	Pharm/Tox telecon scheduled for	Project Manager			
	12/11/02.				
12/4/02	EMail to FDA regarding 12/4/02	Dr. Nitin Patel,	11/22/02	Agency	EMail from FDA with updated
	telecon – outlines status of BI	Project Manager		Contact	information on the 12/4/02 telecon
	minutes and includes word version of	)			
	question list for EOP II Meeting.				
11/22/02	EMail from FDA with updated	Dr. Nitin Patel,	N/A	N/A	N/A
	information on the 12/4/02 telecon.	Project Manager			
11/21/02	Series of e-mails of correspondence	Dr. Nitin Patel,	10/30/02	181	Response to FDA comments - FDA
	involved with rescheduling the	Project Manager			letters dated Aug 29, 2002(Special
	telecon to discuss PK/Tox Issue from				Protocol Assessment, 1182.12) and
	SPA.				Sep 20, 2002 (ddl Drug Interaction)
11/19/02	Rescheduling of FDA 11/20/02	Dr. Nitin Patel,	10/30/02	181	Response to FDA comments - FDA
	PK/Tox Telecon.	Project Manager			letters dated Aug 29, 2002(Special
					Protocol Assessment, 1182.12) and
					Sep 20, 2002 (ddI Drug Interaction)

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	Response to FDA comments – FDA letters dated Aug 29, 2002(Special Protocol Assessment, 1182.12) and Sep 20, 2002 (ddl Drug Interaction)	Response to FDA comments – FDA letters dated Aug 29, 2002(Special Protocol Assessment, 1182.12) and Sep 20, 2002 (ddl Drug Interaction)	Response to FDA comments – FDA letters dated Aug 29, 2002(Special Protocol Assessment, 1182.12) and Sep 20, 2002 (ddl Drug Interaction)	Teleconference held on September 27, 2002	BI's October 11, 2002 letter	EOP2 & F/U on Special Protocol Assessment telecon	Request for Type B Meeting End of Phase IJ/Request for Teleconference Clinical/Pharmacokinetics
ייייייייייייייייייייייייייייייייייייייי	181	181	181	Telecon	177	Agency Contact	176
	10/30/02	10/30/02	10/30/02	9/27/02	10/11/02	10/15/02	10/12/02
NOONT I	Dr. Nitin Patel, Project Manager	Dr. Nitin Patel, Project Manager	Dr. Nitin Patel Project Manager		Dr. Nitin Patel Project Manager		Dr. Nitin Patel, Project Manager
	FDA requests Telecon to discuss drug interaction portion of EOP 2 Package/FDA will send comments from 11/7 internal meeting.	Clarify Issues regarding planning for TPV EOP 2 Meeting and Package.	Mr. Patel requested 5 desk copies of Submission S/N 181 in preparation for a teleconference on 11/20/02.	Final Internal Meeting Minutes for Teleconference held on September 27 discussing Starting Materials in the Synthesis of TPV drug substance.	End of Phase 2 Meeting – Request for Dates Request for Teleconference on BI Responses to Special Protocol	Assessment.	Re: End of Phase 2 Meeting Request and Follow-Up/Special Protocol Assessment Follow-up Teleconference.
	11/8/02	11/13/02	11/1/02	10/23/02	10/22/02		10/15/02

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BI/FDA Teleconference – September 27 <sup>th</sup> – FDA Minutes	BI/FDA Teleconference – September 27th BI Minutes	TPV Special Protocol Assessment 1182.12	Request for Special Protocol Assessment – RESIST 1 Protocol 1182.12
N/A	N/A	Agency Contact	161
9/27/02	9/27/02	8/28/02 & 8/29/02	9/20/02
Dr. Nitin Patel, CSO Dr. Stephen Miller, Team Leader Dr. Dan Boring, Chemistry Reviewer	Dr. Nitin Patel, CSO Dr. Steven Miller, Chemistry Team Leader Dr. Dan Boring, Reviewing Chemist	Mr. Nitin Patel Project Manager	
10/25/02, 10/16/02, BI requested clarification of the minutes provided by FDA by email.	To discuss the starting materials in the synthesis of Tipranavir drug substance.	Re: Tipranavir Special Protocol Assessment 1182.12 Follow-up.	
10/25/02 10/25/02	9/27/02	9/18/02	

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General Correspondence Starting materials	BI notifying of fax sent with the information requested regarding	proposal for 4 TPV as a starting material and to ask for confirmation	of receipt. BI also followed-up on	request for teleconference to discuss 4 TPV as a starting material.	Request for Special Protocol	Assessment 1182.12	Request for desk copies of SPA			Request for Special Protocol	Assessment-Clinical BI requests	special protocol assessment of	1182.12		
141	ACK ACR	ACR			161		ACR			161		ACR			
3/21/02	8/2/02 8/12/02	8/13/02	•		7/18/02		7//19/02			7/18/02		7/30-31/02			
Dr. Dan Boring Reviewing Chemist	DAVDE				Mr. Destry	Sillivan, Project	Manager Ms Virginia	Yoerg, Project	Manager	Mr. Destry	Sillivan,	Project Manager	Ms. Virginia	Yoerg,	Project Manager
To follow-up on setting a date for telecon to discuss starting material designation in the tipranavir drug	substance synthesis.				TPV Special Protocol Assessment					FDA can not approve Request for	Special Protocol Assessment because	submission did not meet	requirements.		
8/30/02					8/28-8/29/02					8/28/02					

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BI notifying of fax sent with the Dan Boring
DAVDP, FDA
Nitin Patel
DAVDP, FDA

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		PERSON		Reference	
FDA email with informal acceptance of ritonavir dose level to be used in rat carcinogenicity study and inquiry about dates of past IND Annual Reports	e >	Nitin Patel DAVDP, FDA	7/19/02	162	Information Amendment- Pharmacology/Toxicology Reports Request for FDA Feedback/Teleconference – BI request concurrence with lowering dose level of ritonavir
			6/7/02 6/18/02 6/19/02 6/25/02 7/15/02 7/24/02	ACR ACR ACR ACR ACR	Request for Feedback on rat carcinogenicity study
FDA called to advise BI of faxed response from Pharm/Tox reviewers regarding acceptance of rat carcinogenicity dose levels and required 3 additional dask coning of	پ يو ا	Nitin Patel DAVDP, FDA	6/7/02 6/18/02 6/19/02 6/25/02	ACR ACR ACR	Feedback on rat carcinogenicity study
the Request for Special Protocol Assessment	-		7/23/02	161	Telephone contact
			7/18/02	161	Request for Special Protocol Assessment-Clinical BI requests special protocol assessment of 1182.12

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7/22/02	FDA emailed with request for	Nitin Patel	7/18/02	191	Request for Special Protocol
7/19/02	additional desk copies of Special	DAVDP, FDA		-	Assessment-Clinical BI requests
	Protocol Assessment				special protocol assessment of 1182.12
7/15/02	BI email FDA advising of a proposed	Nitin Patel	5/24/02	151	Information Amendment:
	decrease in RTV dose in the rat	DAVDP, FDA			Pharmacology/Toxicology report
	carcinogenicity study due to new				and
	Abbott data.				Request for FDA Feedback – BI
					seeking FDA concurrence with dose
					selection for the TPV/RTV and RTV
					groups for 2-year rat carcinogenicity
					study
7/3/02	BI email FDA inquiring about when	Nitin Patel	2/14/02	137	Information Amendment:
	to expect feedback on submissions	DAVDP, FDA			Clinical/Request for Comment-
	regarding the need for ddI drug				Discontinuation of Tipranavir Study
	interaction study				1182.42

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	General Correspondence – BI seeking concurrence with designation of two starting materials in the synthesis of tipranavir drug substance	Request for feedback on starting materials	GC – starting materials GC – Definition of starting materials	Starting Materials	Information Amendment:	Clinical/Request for Comment- Discontinuation of Tipranavir	Study 1182.42	Information Amendment: Pharmacology/Toxicology report	and	Request for FDA Feedback - BI	seeking FDA concurrence with dose selection for the TPV/RTV	and RTV groups for 2-year rat carcinogenicity study
Reference	141	ACR	095 106	ACR	137			151				
	3/21/02	8/28/01	6/6/01 8/22/01	8/22/01	2/14/02			5/24/02				
PERSON	Dan Boring DAVDP, FDA				Nitin Patel	DAVDP, FDA						
	FDA called BI in regard to their review of BI's proposal regarding the starting materials				BI emailed and phoned FDA about	the need for FDA's feedback on the rat carcinogenicity study and the need	for conducting another ddI drug interaction study. FDA faxed	response to rat carcinogenicity study submissions but acknowledged that	the other response was pending on	the biopharm reviewer.		
	6/17/02				6/7/02	6/18/02 6/19/02	6/25/02					

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	TO CONTRA		NOICE CHICA	
BI emailed FDA seeking the status of Nitin Patel FDA's feedback regarding need for DAVDP, F ddI Drug Interaction study.	Nitin Patel DAVDP, FDA	2/14/02	137	Information Amendment: Clinical/Request for Comment- Discontinuation of Tipranavir Study 1182.42
يو	Nitin Patel DAVDP, FDA	4/26/02	146	Information Amendment – Clinical Investigators Brochure Version 5
BI called FDA to confirm receipt of submission that was faxed and to reiterate the request for feedback on the Phase II Protocol (1182.52)	Karen Young DAVDP, FDA	3/5/02	139	Response to FDA Request for Information
FDA informed BI of an internal attempts to set up teleconference with BI to discuss formulation and Pharmacology/Toxicology questions		N/A	N/A	N/A
BI's minutes from teleconference between BI and FDA	Karen Young DAVDP, FDA & TPV Team	2/15/02	N/A	Fax from FDA Requesting Information from BI in order to address the clinical development program. FDA Requesting Chemistry, Pharmacology/Toxicology, Pharmacokinetics and Clinical information.

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Information Amendment - Clinical	Pharmacokinetics from 1182.6			DI Feedback					Information Amendment - Clinical	Pharmacokinetics from 1182.6						Information Amendment – CMC	support for on-going and future	clinical trails		
132				ACR				**	132			ACR	Contact			100				
12/20/01				1/29/02					12/20/01			1/29/02				6/19/01				
Karen Young	DAVDP, FDA								Karen Young	DAVDP, FDA						Karen Young	DAVDP, FDA			
FDA called notifying BI that the	Division will be having an internal	meeting today to discuss the current	status of TPV and what might be	needed to support Phase III. FDA	said they'd be sending	pharmacokinetic, chemistry,	pharm/tox and clinical comments in	the next week.	BI called FDA notifying them of the	email of the submission on the	tipranavir formulation, to inquire	about the need for a teleconference	and requesting feedback on the drug	interaction program in support of	Phase 3.	FDA called requesting more	information on SEDDS formulation	used in clinical trials. BI responded	saying that more information will be	,
2/07/02						-			1/29/02							1/29/02				

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Response to FDA Comments in regards to combination toxicology studies	Information Amendment - Clinical Pharmacokinetics from 1182.6	Comment on 1182.6	Telephone contact			Telephone contact	•		Type C Meeting	
133	132	ACR	N/A			N/A			N/A	Agency Contact
12/27/01	12/20/01	9/25/01	N/A			N/A			10/5/01	11/7/01
Karen Young DAVDP, FDA			Karen Young DAVDP, FDA			Karen Young	DAVDP, FDA		Leslie Stevens	DAVDP, FDA
FDA called to request additional information on the combination tox studies. BI asked FDA about data from 1182.6 study and requested	comments on whether or not data was sufficient.		BI	ddI which was being conducted in Germany. BI ensured FDA that	background information regarding	FDA called BI to	<ul> <li>Advise project manager has changed for tipranavir</li> </ul>	<ul> <li>Discuss and request information on the SEDDS formulation</li> </ul>	BI called FDA to check on the status	of FDA's Meeting Minutes from Type C Meeting
1/28/02			12/13/01 12/18/01			12/12/01	12/13/01		11/30/01	

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11/15/01	BI called FDA to follow-up on status	Leslie Stephens	11/2/01	122	Request for Advice on Duration of
	of BI's Request for Advice on Duration of Combination Toxicology Studies	Project Manager DAVDP			Combination Toxicology Studies
11/07/01	BI called FDA to:	Leslie Stephens	10/5/01	N/A	Type C Meeting
	Advise Division of fax coming	Project Manager			
	containing BI's proposal for	DAVDP			
	combination toxicology studies				4
	Advise Division that BI's meeting				
	minutes from Type C meeting will				
	also be faxed				
10/26/01	BI called FDA to:	Leslie Stephens	10/5/01	N/A	Type C Meeting
	Advise that BI plans on submitting	Project Manager			
	a proposal for combination	DAVDP			
	toxicology studies and request				
	discussion of timing of				
	submission, review and feedback				
	of proposal				
	Follow-up on Division's offer to				
	provide a draft template for				
	genotyping and phenotyping				
9/26/01	BI called FDA to confirm extent of	KarenYoung	9/25/01	115	Desk Copies of General
	the background document that was	DAVDP, FDA			Correspondence - Background
	needed electronically.				Document for Type C Meeting

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9/25/01	FDA called to confirm internal	KarenYoung	3/16/01	082	Response to FDA comments faxed
	meeting on 10/1/01 and give feedback on Protocol 1182.6	DAVDP, FDA		}	on March 3 – 5, 2001 regarding
9/24-25/01	FDA called questioning the eligibility	Leslie Stephens	1/19/01	920	Protocol Amendment - New
	criteria regarding how patients were	Project Manager			Protocol and New Investigator for
	assigned to the nucleosides/non-	DAVDP			1182.6
	nucleosides and requested two	KarenYoung			
	additional copies of the background	DAVDP, FDA	9/25/01	115	Desk Copies of General
	document				Correspondence - Background
					Document for Type C Meeting
8/28/01	FDA informed BI of their response to		10/9/9	560	General Correspondence –
	the proposal for 5 TPV as a starting	DAVDP, FDA			requesting FDA concurrence with
	material in the synthesis of tipranavir				designation of starting materials in
	drug substance. Bi asked for				the syntheses of tipranavir drug
	feedback on the original proposal.				substance
			8/22/01	901	General Correspondence - Proposal
					for the definition of starting
					materials in synthesis of TPV drug
					substance
			7/18/01	N/A	ACR - FDA called BI regarding the
					proposal for starting materials

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8/24/01	BI called FDA to confirm BI's availability for Type C meeting scheduled for October 5, 2001	Leslie Stephens Project Manager DAVDP	6/21/01	ACR	Meeting Plans
8/22/01	BI called FDA to:  Obtain status update on review of starting material proposal  Ask how FDA's new position on starting materials would be nublished	Dan Boring DAVDP, FDA	6/6/01	095	General Correspondence – requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance
	Obtain clarification on one of the existing (old) criteria for starting materials     Inquire about future CMC IND amendments		8/22/01	901	General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance
7/18/01	FDA called BI regarding the proposal for starting materials	Dan Boring DAVDP, FDA	6/6/01	960	General Correspondence – requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance
6/21/01	FDA called with date/time proposal for Type C Meeting teleconference.	Destry Sullivan DAVDP, FDA	6/20/02	Telecon	Telephone contact

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6/20/01	CO	Destry Sullivan	4/5/01	N/A	Pre-IND/End of Phase I Meeting
	Meeting Request. FDA wanted to	DAVDP, FDA			
	propose a teleconference rather than a				
	face-to-face meeting. BI explained				
	reasoning for face-to-face meeting				
	request and FDA said they'd get back				
	to her on it. BI also inquired about				
	feedback on safety update.				
6/08/01	BI called FDA to:	Anthony DeCicco	4/5/01	N/A	Pre-IND/End of Phase I Meeting
	Request a meeting in late July to	DAVDP, FDA			)
	discuss plans for the TPV pivotal				Request for safety data & feedback
	program		4/3/01	ACR	on DI proposal
	Request feedback on availability				
	of meeting minutes from April 5,				
	2001				
	<ul> <li>Request feedback on requested</li> </ul>				
	safety update				
4/3/01	FDA called in regards to the	Leslie Stephens	N/A	N/A	Telephone contact
	upcoming pre-NDA Meeting. FDA	Project Manager			
	requested that safety data be provided	DAVDP			
	and that BI be ready to discuss drug	Joseph Toerner,			
	interaction studies for tipranavir.	DAVDP, FDA			

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		PERSON		Keference	
2/8/01	BI call FDA to confirm End of Phase I meeting date for April 5, 2001	Leslie Stephens Project Manager DAVDP	1/31/01	N/A	FDA requested change of date for the End of Phase 1 meeting date
		, .	2/6/01	N/A	BI Request change to upcoming End of Phase 1 meeting date
			2/7/01	N/A	BI and FDA discuss potential meeting dates for Tipranavir End of Phase I meeting
2/7/01	BI and FDA discuss potential meeting dates for Tipranavir End of Phase I meeting	Leslie Stephens Project Manager DAVDP	1/31/01	N/A	FDA requested change of date for the End of Phase 1 meeting date
. ,			2/6/01	N/A	BI Request change to upcoming End of Phase 1 meeting date
2/6/01	BI Request change to upcoming End of Phase 1 meeting date	Leslie Stephens Project Manager DAVDP	1/31/01	N/A	FDA requested change of date for the End of Phase 1 meeting date
1/31/01	FDA requested change of date for the Leslie Stephens End of Phase 1 meeting date DAVDP	Leslie Stephens Project Manager DAVDP	N/A	N/A	N/A
1/15/01	BI called to confirm BI's meeting date of 3/1/01.	Leslie Stephens Project Manager DAVDP	N/A	N/A	N/A

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Protocol Amendment – Change in Protocol M/3342/0006 Amendment 2	Telephone contact	Protocol Amendment —  New Protocol M/3342/0013  Change in Protocol M/3342/0006 Amendment 1  New Investigators for Protocol M/3342/0006	FDA Request for information – Clinical Comments on SN 040 and SN 041	FDA Request for information – Clinical Comments on SN 040 and SN 041
053	N/A	043	040 041	0 <del>4</del> 0 041
8/20/99	86/6	3/29/04	3/24/99	3/24/99
Leslie Stevens DAVDP, FDA	Christine Kelly DAVDP, FDA	Christine Kelly Joe Toerner DAVDP, FDA	Christine Kelly Joe Toerner DAVDP, FDA	Christine Kelly DAVDP, FDA
FDA called with questions regarding Protocol M/3342/0006. PNU ensured they would look into them and get back as soon as possible.	PNU called to clarify if submitting 16 weeks data for naïve patients instead of 24 weeks when filing the NDA for accelerated approval would be acceptable. FDA said they would have to wait until the Phase II studies are complete to see what the data looks like.	FDA called to inform PNU that protocols (13 and 19) were missing from SN 043. PNU told FDA that they would resubmit SN 043.	FDA called to remind PNU that 26-week rat and 39-week dog reports. Were due by April 2, 1999 or clinical study M/3342/0006 would be put on clinical hold.	Call made regarding request for animal toxicology reports.
9/16/99	5/3/99	4/6/99	3/29/99	3/24/99-3/23/99

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9/22/98	PNU notified the FDA about a minor	David Staten	9/23/98	032	Protocol Amendment -New
	error on cover sheet on Protocol	DAVDP, FDA			Protocol M/3342/0011 - Draft
	M/3342/0011. "Final" protocol				Information Amendment – CMC
	should have been stated rather than				data including new SEDDs
	"Draft". FDA asked for new cover				formulations and Toxicology Report
	page be sent.				a0014475
9/21/98	FDA called notifying PNU that	Debra Gump	3/18/98	021	Information Amendment - CMC
	"boiler plate" investigational drug	DAVDP, FDA			information on the synthesis of drug
	statement was missing from final				substance, new formulation, draft
	labels that were being used in studies.				labeling, and changed storage
	PNU ensured correction to be sent.				condition for 150 mg capsules
86/8/6	FDA called to reschedule Sept 9	Debra Gump	N/A	N/A	N/A
	teleconference due to conflict in	DAVDP, FDA	,		
	attendee's schedule. PNU decided to	-			
100	proceed without that individual.	-			
	FDA request more information on the				
	subject of discussion.				
8/21/98	Contact regarding the background	Debra Gump	N/A	N/A	N/A
	document for the September 8th	DAVDP, FDA			
	teleconference between PNU and				
	FDA				

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7/29/98	PNU call FDA to confirm September	Debra Gump	7/28/98	N/A	Telephone contact
	9th is ok with the PNU team for the	DAVDP, FDA			
	teleconference to discuss further				
	development plans for tipranavir.				
	PNU promises background				
	information to be sent shortly.				
7/28/98	PNU call FDA to confirm September	Debra Gump	N/A	N/A	N/A
	date for teleconference to discuss	DAVDP, FDA			
	further development plans for				
	tipranavir. FDA offered a Sept. 9				
	date.				
86/11/9	FDA approved changes to Protocol	Debra Gump	86/11/98	N/A	PNU fax containing proposed
	M/3342/0004 that were previously	DAVDP, FDA			changes to Protocol M/3342/0004
	ומאכט וט ווכן				
					Protocol Amendment - Change in
			86/7/9	025	Protocol M/3342/004 Amendment 8
					Amendment A, 6 and 7

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		CONTACT	Reference	of Reference	•
9/11/98	Called FDA to confirm receipt of fax with proposed amendment to Protocol M/3342/004	Debra Gump DAVDP, FDA	86/2/98	025	Protocol Amendment – Change in Protocol M/3342/004 Amendment 8
2/17/98	FDA relayed message that protocol M/3342/0004 (Amendment 9) were ok with Division.				
2/11/98	Call placed to FDA to confirm fax of proposed amendment to Protocol M3342/0004. It was confirmed.				
9/10/97	Inform FDA of plans of submitting 3-month interim reports for each 6-month animal study	Kimberly A. Struble DAVDP, FDA	26/2/9	N/A	Telephone contact
	•	,	5/15/97	N/A	Request for feedback to the strategy of conducting 6-month instead of 3-month toxicity study
			4/15/97	900	Information Amendment: Pharmacology/ Toxicology - Update to Gantt chart of proposed clinical trails together with one for proposed preclinical studies designed to support the safety, Final Toxicology Reports
6/5/97	FDA Request to prepare to submit for Kimberly A.	Kimberly A.	5/15/97	N/A	Request for feedback to the strategy

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		CONTACT PERSON	Reference	of Reference	
					Consider a second secon
	each 6 month animal study a 6 month interim report	Struble DAVDP, FDA			of conducting 6-month instead of 3-month toxicity study
	•				
			4/15/07	900	Information Amendment:
			1001	000	Update to Gantt chart of proposed
					proposed preclinical studies
					designed to support the safety, Final Toxicology Reports
96/9/9	PN&U requested FDA opinion on	Kimberly A.	N/A	N/A	N/A
	dose level selection for the 30 day	Struble			
	dog study for tipranavir. FDA agreed	DAVDP, FDA			
	on the 320 mg/kg/day.				
11/20/96	IND assigned 51-979	Dawn M. Roberts	N/A	N/A	References for IND for PNU- 140690
12/16/96	FDA telephone contact on 12/13/96	Dawn M. Roberts	N/A	N/A	December 13, 1996 Safety Review
	concerning PNU-140690 for AIDS,				meeting - FDA determined it is safe
	AIDS related complex and treatment				for P&U to proceed with the single
	of Asymptomatic HIV Positives				oral dose escalation study in healthy
	RESCRIPTOR Tablets				volunteers (protocol M/3342/0001)
12/13/96	FDA determined it was safe for P&U	Kimberly A.	12/13/96	100	Protocol Amendment - Change in
	to proceed with the single oral dose	Struble			Protocol M/3342/001 - Amendment
	escalation study in healthy	DAVDP, FDA			
	volunteers. (Protocol M/3342/0001)				

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1	Description of Submission	Date of	Serial No.	Description	Date of Reference
Original IND Application	Application	11/13/96	N/A	ot Reference N/A	
Protocol Amer Protocol M/33 IRB Letter of , Consent Form of Protocol M/ Methodist Hos Committee	Protocol Amendment - Change in Protocol M/3342/001 - Amendment 1 IRB Letter of Approval and Patient Consent Form for evidence of approval of Protocol M/3342/0001 by the Bronson Methodist Hospital Human Use Committee	12/13/96	000	Original Protocol	11/13/96
Protocol Amer Protocol M/33	Protocol Amendment – Change in Protocol M/3342/001 Amendment 2	1/1/97	000	Original Protocol	11/13/96
		:	001	Amendment 1 to M/3342/0021	12/13/96
Protocol Amer M/3342/002 (( information) Information A Report	Protocol Amendment - New Protocol M/3342/002 (CMC and labeling information) Information Amendment - Toxicology Report	1/2/97	N/A	N/A	N/A
General Corresponder FDA faxes containing Pharmacology, Micro Chemistry comments	General Correspondence - Response to FDA faxes containing Clinical, Pharmacology, Microbiology, and Chemistry comments	1/31/97	N/A	FDA faxes	12/17/96 12/19/96

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
S N	$\downarrow$		Submission	Referenced	of Reference	
900	•	Protocol Amendment - Change in Protocol M/3342/002 Amendment 1	2/20/97	001	Original Protocol	12/13/96
	•	Information Amendment - Toxicology Reports				
	•	Correction to General Correspondence - SN 004 submitted 1/31/97				
900	•	Information Amendment: Pharmacology/ Toxicology - Update to Gantt chart of	4/15/97	004	Gantt chart submitted	1/31/97
		proposed clinical trails together with one		000	Draft Toxicology reports	11/13/96
		for proposed preclinical studies designed to support the safety				
	•	Final Toxicology Reports,				
200	•	Information Amendment: Toxicology	26/1/97	N/A	N/A	N/A
	_	Keport				
800	•	Protocol Amendment - New Protocol	<i>L6/6/S</i>	N/A	N/A	N/A
		M/3342/0003 and New Investigator, Dr. Dennis W Schneck				
	•	Information Amendment –				
		• CMC				
		<ul> <li>Updated stability data</li> </ul>				
		<ul> <li>Labeling for Protocol M/33420003</li> </ul>				
600	•	Protocol Amendment –	7/3/97	N/A	N/A	N/A
		New Protocol and Investigator for				
		MI/3342/0005				

Description of Submission         Date of formation Amendment – Labeling for M/3342/0004         Serial No. of Reference         Description of Reference           • Information Amendment – Labeling for M/3342/0004         • Protocol Amendment – New Protocol Amendment – New Investigator, for M/3342/0004         8/4/97         N/A         N/A         N/A           • Protocol Amendment – New Investigator, for M/3342/0004         • Information Amendment – New Investigator, for M/3342/0004         8/4/97         010         Original protocol         7/1           • CMC information Amendment – Information an support of the trial product         • Labeling for M/3342/0004         8/22/97         010         Original Protocol         7/1           • Protocol Amendment – Toxicology Reports         Information Amendment – Toxicology Reports         8/22/97         010         Original Protocol         7/1           • General Correspondence – P&U Minntes         9/25/97         N/A         Teleconference between P&U and proposal to extend the ongoing 26-week         Protocol Amendment – Change in Am					Q	
• Information Amendment – Labeling for M/3342/0005         Submission Referenced         Referenced of Reference           • Protocol Amendment – New Protocol Amendment – New Investigator, for M/3342/004         8/4/97         N/A         N/A           • Protocol Amendment – New Investigator, for M/3342/004         8/4/97         010         Original protocol           • Information Amendment – Change in Information Amendment – Change in Information Amendment – Toxicology/Pharmacology Reports         8/22/97         010         Original Protocol           • Information Amendment – Toxicology Support for Protocol M/3342/0004.         8/27/97         010         Original Protocol           • Information Amendment – Toxicology Support for Protocol M/3342/0004.         8/25/97         N/A         Teleconference between P&U and protocol           • Information Amendment – Change in Protocol Amendment 2         10/10/97         010         Original Protocol Amendment 1           • Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2         10/10/97         010         Original Amendment 1           • Info Amend – Tox.Rep. TR 7228-97-052         010         Original Amendment 1	Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
• Information Amendment – Labeling for M/3342/0005         7/10/97         N/A         N/A           • Protocol Amendment – New Protocol Amendment – New Investigator, for M/3342/004         8/4/97         010         Original protocol           • CMC information in support of the trial product         • Labeling for M/3343/004         8/22/97         010         Original Protocol           • Protocol Amendment – Change in Protocol M/3342/0004 Amendment – Toxicology/Pharmacology Reports         8/22/97         010         Original Protocol           • Information Amendment – Toxicology Pharmacology Reports         8/22/97         010         Original Protocol           • Information Amendment – Toxicology Pharmacology Reports         8/22/97         010         Original Protocol           • Information Amendment – Toxicology Support for Protocol M/3342/0004.         8/22/97         010         Original Protocol           • General Correspondence – P&U Minutes of Teleconference that discussed P&U's proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks non-rodent Amendment 2 non-rodent Tox.Rep. TR 7228-97-052         010         Original Amendment 1 non-rodent Amendment 2 non-rodent Amendment	% V			Referenced	of Reference	
<ul> <li>Protocol Amendment - New Protocol</li></ul>		Information Amendment – Labelin, for M/3342/0005				
<ul> <li>Protocol Amendment – New Investigator, for M/3342/004</li> <li>Information Amendment – New Investigator, for M/3342/004</li> <li>CMC information in support of the trial product</li> <li>Labeling for M/3343/004</li> <li>Protocol Amendment – Change in Protocol M/3342/0004.</li> <li>Information Amendment – Toxicology Reports</li> <li>Information Amendment – Toxicology Reports</li> <li>Information Amendment – Toxicology Support for Protocol M/3342/0004.</li> <li>General Correspondence – P&amp;U Minutes of Teleconference that discussed P&amp;U's proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks non-rodent toxicology study to 39 weeks non-rodent toxicology study to 39 weeks and 3</li> <li>Protocol M/3342/0004 – Amendment 2 and 3</li> <li>Info Amend – Tox.Rep. TR 7228-97-052</li> </ul>	010	Protocol Amendment – New Protocol M/3342/004	7/10/97	N/A	N/A	N/A
<ul> <li>Information Amendment -</li> <li>CMC information in support of the trial product</li> <li>Labeling for M/3343/004</li> <li>Protocol Amendment - Change in Protocol M/3342/0004 Amendment 1</li> <li>Information Amendment - Toxicology/Pharmacology Reports</li> <li>Information Amendment - Toxicology/Pharmacology Reports</li> <li>Information Amendment - Toxicology S/27/97</li> <li>Information Amendment - Toxicology S/25/97</li> <li>Information Amendment - P&amp;U Minutes</li> <li>General Correspondence - P&amp;U Minutes</li> <li>General Correspondence - P&amp;U Minutes</li> <li>Greleconference that discussed P&amp;U's proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks</li> <li>Protocol Amendment - Change in Protocol M/3342/0004 - Amendment 2 and 3</li> <li>Info Amend - Tox.Rep, TR 7228-97-052</li> </ul>	011	<ul> <li>Protocol Amendment – New Investigat for M/3342/004</li> </ul>		010	Original protocol	7/10/97
• Labeling for M/3343/004         # Protocol Amendment – Change in Protocol Amendment – Change in Protocol M/3342/0004 Amendment – Toxicology/Pharmacology Reports         8/22/97         010         Original Protocol           • Information Amendment – Toxicology/Pharmacology Reports         8/27/97         010         Original Protocol           • Information Amendment – Toxicology Pharmacology Reports         8/27/97         010         Original Protocol           • Information Amendment – Toxicology Support for Protocol M/3342/0004.         9/25/97         N/A         Teleconference between P&U and Protocol           • General Correspondence – P&U Minutes of Teleconference that discussed P&U's proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks         9/25/97         N/A         Teleconference between P&U and Protocol M/3342/0004 – Amendment 2           • Protocol M/3342/0004 – Amendment 2 and 3         010         Original Protocol M/3342/0004 – Amendment 2           • Info Amend – Tox.Rep, TR 7228-97-052         012         Amendment 1		upport				
<ul> <li>Protocol Amendment – Change in Protocol M/3342/0004 Amendment 1  - Information Amendment – Toxicology/Pharmacology Reports - Information Amendment – Toxicology/Pharmacology Reports - Information Amendment – Toxicology/Pharmacology Reports - Information Amendment – Toxicology Reports - Info Amend – Toxicology Reports - Information Amendment – Change in Protocol M/3342/0004 – Amendment 2 - Information Amendment I Amendment I Amendment I - Info Amend – Toxicology Reports - Info Amend – Toxicolog</li></ul>		trial product  • Labeling for M/3343/004				
<ul> <li>Information Amendment –         Toxicology/Pharmacology Reports     </li> <li>Information Amendment – Toxicology         Support for Protocol M/3342/0004.     </li> <li>General Correspondence – P&amp;U Minutes of Teleconference between P&amp;U and proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks         Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2 and 3     </li> <li>Info Amend – Tox.Rep, TR 7228-97-052</li> </ul>	012	<ul> <li>Protocol Amendment – Change in Protocol M/3342/0004 Amendment 1</li> </ul>	8/22/97	010	Original Protocol	7/10/97
<ul> <li>Information Amendment – Toxicology Support for Protocol M/3342/0004.</li> <li>General Correspondence – P&amp;U Minutes of Teleconference that discussed P&amp;U's proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks non-rodent toxicology study to 39 weeks and 3</li> <li>Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2 and 3</li> <li>Info Amend – Tox.Rep, TR 7228-97-052</li> </ul>		Information Amendment –     Toxicology/Pharmacology Reports				
<ul> <li>General Correspondence – P&amp;U Minutes of Teleconference between P&amp;U and of Teleconference that discussed P&amp;U's proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks</li> <li>Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2 and 3</li> <li>Info Amend – Tox. Rep, TR 7228-97-052</li> </ul>	013	<ul> <li>Information Amendment – Toxicology Support for Protocol M/3342/0004.</li> </ul>	8/27/97	010	Original Protocol	7/10/97
<ul> <li>proposal to extend the ongoing 26-week non-rodent toxicology study to 39 weeks</li> <li>Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2 and 3</li> <li>Info Amend – Tox.Rep, TR 7228-97-052</li> </ul>	014	• General Correspondence – P&U Minut of Teleconference that discussed P&U'		N/A	Teleconference between P&U and FDA on September 22, 1997	9/22/97
<ul> <li>Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2 and 3</li> <li>Info Amend – Tox. Rep, TR 7228-97-052</li> </ul>		proposal to extend the ongoing 26-wee non-rodent toxicology study to 39 weel	<u>s</u>			
• Info Amend – Tox.Rep, TR 7228-97-052	015	<ul> <li>Protocol Amendment – Change in Protocol M/3342/0004 – Amendment 2 and 3</li> </ul>		010 012	Original Amendment 1	7/10/97 8/22/97
		• Info Amend - Tox.Rep, TR 7228-97-0	1.5			

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
0	-		Submission	Keterenced	of Keterence	
016	•	Protocol Amendment – New Protocol and Investigator for M/3342/0007	10/24/97	N/A	N/A	N/A
	•	Information Amendment - CMC				
		information and labeling in support of M/3342/0007				
017	•	Protocol Amendment - New Protocol	12/23/97			
		and Investigator for M/3342/008		4/14		
	•	Amendment of IND to reflect proposal to discontinue the routine performance of		N/A	the routine performance of thyroid	11/10/9/
		thyroid function testing in new clinical			function testing in new clinical	
		trials with PNU-140690			trials on 11/10/97	
	•	Information Amendment		N/A		11/14/97
	•	Labeling for M/3342/0008			Approval of proposal	
	•	Pharmacology /Toxicology Reports			communication by FDA	
018	•	Protocol Amendment –	2/4/98			
		New Protocol DRAFT M/3342/0009		N/A	N/A	N/A
		Change in Protocol –		010	Original Protocol	7/10/97
		M/3342/0004 Amendments 4 and 5		012	Amendment 1	8/22/97
				015	Amendments 2 and 3	10/10/97
		• M/3342/0008 Amendments 1		017	Original Protocol	
	_					12/23/97

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
	and 2				
-	1000/0700/14 3 1 1 1 1 1 1 1		010	Original Protocol	7/10/97
	Inew investigator for MJ/3542/0004		012 015	Amendments 1 Amendments 2 and 3	8/22/97 10/10/97
	Information Amendment: Pharmacology and ADME Reports				
010	<ul> <li>IND Annual Report - reporting period through 9/30/97</li> </ul>	2/6/98	N/A	N/A	N/A
020	<ul> <li>Protocol Amendment – FINAL Protocol and New Investigator M/3342/0009</li> </ul>	2/25/98	018	Draft Protocol M/3342/0009	2/4/98
_	• Labeling in support of Protocol M/3342/0009				
	Information Amendment –     Pharmacology/Toxicology				
021	Information Amendment – CMC information on the synthesis of drug	3/18/98	800	Information Amendment - CMC	5/9/97
	substance, new formulation, draft labeling, and changed storage condition for 150 mg conculas				
	tot too till capsures				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
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022	<ul> <li>Protocol Amendment – Change in</li> </ul>	4/1/98	010	Original Protocol	7/10/97
	Protocol M/3342/0004 Amendments A,		012	Amendment 1	8/22/97
	6, and 7		015	Amendment 2 and 3	9/30/97
			018	Amendment 4 and 5	2/4/98
023	Protocol Amendment – Change in	4/30/98	018	Draft protocol	2/4/98
	Protocol M/3342/0009 Amendment 1		020	Final Original Protocol	2/25/98
024	<ul> <li>Protocol Amendment – New</li> </ul>	8/17/9	010	Original Protocol	7/10/97
	Investigators for Protocol M/3342/0004		012	Amendment 1	8/22/97
			015	Amendment 2 and 3	9/30/97
			018	Amendment 4 and 5	2/4/98
			022	Amendment 6 and 7	4/1/98
025	Protocol Amendment – Change in	6/2/98	010	Original Protocol	7/10/97
	Protocol M/3342/004 Amendment 8		012	Amendment 1	8/22/97
			015	Amendment 2 and 3	9/30/97
			018	Amendment 4 and 5	2/4/98
			022	Amendment 6 and 7	4/1/98
970	General Correspondence – Meeting	7/13/98	N/A	N/A	N/A
	Request to review BI's pharmacokinetic, efficacy and safety data				
027	<ul> <li>Protocol Amendment – Change in</li> </ul>	2/30/98	010	Original Protocol	7/10/97
	Protocol M/3342/0004 Amendment 9		012	Amendment 1	8/22/97
			015	Amendment 2 and 3	9/30/97

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
OK		Submission	Kererenced	of Keference	
			018	Amendment 4 and 5	2/4/98
			022	Amendment 6 and 7	4/1/98
			025	Amendment 8	86/2/98
028	Protocol Amendment – New protocol M/3342/0012	8/25/98	N/A	N/A	N/A
029	General Correspondence –	8/56/98	026	General Correspondence –	7/13/98
	Teleconference Background Package for			Meeting Request	
	September 9, 1998 teleconference to				
	discuss BI's pharmacokinetic, efficacy and safety data				
030	General Correspondence –	9/18/98	026	General Correspondence –	7/13/98
	Teleconference Meeting Minutes from			Meeting Request	
	September 9, 1998 teleconference				
	discussing BI's pharmacokinetic, efficacy		029	General Correspondence –	8/56/98
	and safety data			Teleconference Background Package	
				ò	86/6/6
			N/A	Teleconference between BI and FDA	
031	Information Amendment – Clinical Study Report and Toxicology Report	9/21/98	N/A	N/A	N/A
	Treport alla Loricology treport				
032	Protocol Amendment –New Protocol     M/3342/0011 – Draft	9/23/98	N/A	N/A	N/A
	Information Amendment – CMC data		;		

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S <sub>o</sub>		Submission	Referenced	of Reference	
	including new SEDDs formulations and Toxicology Report a0014475				
033	<ul> <li>General Correspondence – Update to Serial No. 021, new labeling</li> </ul>		N/A	Telephone call between BI and FDA	9/21/98
			021	Information Amendment	3/18/98
034	<ul> <li>General Correspondence – Correction to Serial No. 032 - Final Protocol</li> </ul>	10/5/98	N/A	Telephone call between BI and FDA	9/22/98
	M/3342/0011		032		9/23/98
				Protocol Amendment –New Protocol M/3342/0011 –	
				Draft	
035	General Correspondence – Response to FDA Fax – Biopharmaceutics Comments on Protocol M/3342/0012	10/6/98	028	FDA Fax with Biopharmaceutics comments	9/23/98
036	Protocol Amendment – Change in Protocol M/3342/0012 Amendment 1	10/22/98	028	Original Protocol	825/98
037	Protocol Amendment	11/2/98	N/A	N/A	N/A
	<ul> <li>New Protocol and Investigator for M/3342/0014</li> </ul>				
	Information Amendment				
	Clinical Analysis Memo				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
	<ul><li>Clinical Study Reports</li><li>CMC labeling for M/3342/0004</li></ul>				
	<ul> <li>Pharmacology/Toxicology Reports</li> </ul>				
038	Information Amendment –	12/4/98	N/A	N/A	N/A
	Pharmacology/Toxicology Reports	:			
039	Protocol Amendment –	12/9/98			
	<ul> <li>New Protocol – M/3342/0115</li> </ul>				
	• Changes in Protocol M/3342/0004		010	Original Protocol	1/10/97
	Amendment 10		012	Amendment 1	8/22/97
			015	Amendment 2 and 3	9/30/97
			018	Amendment 4 and 5	2/4/98
			022	Amendment 6 and 7	4/1/98
			025	Amendment 8	9/2/98
			027	Amendment 9	7/30/98
	<ul> <li>Amendment to Clinical Study Report a0024338</li> </ul>		037	Final Trial Report M/3342/0007	11/23/98
040	<ul> <li>Annual Report – Reporting Period 10/1/97 – 9/30-98</li> </ul>	1/29/99	N/A	N/A	N/A
041	<ul> <li>Protocol Amendment – New Protocol M3342/0006</li> </ul>	2/5/99	N/A	N/A	N/A
042	Protocol Amendment – New Investigators for M/3342/0015	3/1/99	039	Original Protocol	12/9/98

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>			Submission	Referenced	of Reference	
043	•	Protocol Amendment –  New Protocol M/3342/0013	3/29/99			
		Change in Protocol M/3342/0006     Amendment 1		041	Original Protocol	2/5/99
		<ul> <li>New Investigators for Protocol M/3342/0006</li> </ul>				
044	•	Information Amendment – CMC, updated sections of drug protocol	3/31/99	000	Original IND Application	11/13/96
045	•	Protocol Amendment - Toxicology Study Reports	4/1/99	N/A	N/A	N/A
046	•	General Correspondence - Response to	4/2/99		FDA Fax	3/24/99
	<del></del>	FDA Questions regarding Submissions SN 040 and 041		040	Annual Report – Reporting Period 10/1/97 – 9/30-98	1/29/99
,				041	Annual Report – Reporting Period 10/1/97 – 9/30-98	2/5/99
047	•	Protocol Amendment – Toxicology Study Report resubmission	4/5/99	045	Protocol Amendment – Toxicology Study Reports a0011536	4/1/99
048	•	General Correspondence – Request for Review of Carcinogenicity Study Protocol	4/6/99	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
049	<ul><li>Protocol Amendment –</li><li>New Protocol M/3342/0016</li></ul>	66/01/5			
	<ul> <li>New Investigator for Protocol M/3342/0006</li> </ul>		041 043	Original Protocol Amendment 1	2/5/99 3/29/99
	<ul> <li>New Investigator for Protocol M/3342/0013</li> </ul>		043	Original Protocol	3/29/99
	<ul> <li>New Investigator for Protocol 69INF0013-019</li> </ul>		043	Original Protocol	3/29/99
	<ul> <li>Change in Investigator for Protocol M/3342/0015</li> </ul>	-00	039	Original Protocol	12/9/98
020	<ul> <li>Protocol Amendment –</li> <li>New Investigator for Protocol M/3342/0006</li> </ul>	5/27/99	041	Original Protocol Amendment 1	2/5/99 3/29/99
	<ul> <li>Information Amendment –Toxicology</li> <li>Data</li> </ul>		N/A	N/A	N/A
051	Protocol Amendment – New Investigators for Protocol M/3342/0006	6/22/99	041	Original Protocol Amendment 1	2/5/99 3/29/99
052	<ul> <li>Protocol Amendment – New Investigators for M/3342/0006</li> </ul>	66/6/8	041 043	Original Protocol Amendment 1	2/5/99 3/29/99

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Š		Submission	Referenced	of Reference	
	Information Amendment –     Pharmacology/Toxicology Data –		N/A	N/A	N/A
053	<ul> <li>Protocol Amendment – Change in protocol M/3342/0006 Amendment 2</li> </ul>	8/20/99	041 043	Original Protocol Amendment 1	2/5/99 3/29/99
054	<ul> <li>Protocol Amendment –</li> <li>Change in Protocol M/3342/0016 –</li> <li>Amendment 1</li> </ul>	6/11/6	049	Original protocol	5/10/99
	<ul><li>New Investigator M/3342/0006</li><li>Added sub-investigators M/3342/0006</li></ul>		041 043 053	Original Protocol Amendment 1 Amendment 2	2/5/99 3/29/99 8/20/99
	<ul> <li>Information Amendment – Clinical Data a0038615</li> </ul>		020	Original Protocol	2/24/98
055	<ul> <li>Desk Copy of Serial No. 044</li> </ul>	9/24/99	044	Information Amendment – CMC updated sections of drug protocol	4/1/99
950	<ul> <li>Desk copy of Serial No. 040</li> </ul>	66/30/6	045	Annual Report – Reporting Period 10/1/97 – 9/30-98	1/29/99
057	<ul> <li>Protocol Amendment – New investigator and added sub-investigator for M/3342/0006</li> </ul>	10/21/99	041 043 053	Original Protocol Amendment 1 Amendment 2	2/5/99 3/29/99 8/20/99
	• Info Amend – Pharm Data a0056253				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
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058	<ul> <li>Information Amendment –     Pharmacology/Toxicology Data</li> </ul>	10/27/99	N/A	N/A	N/A
029	<ul> <li>Protocol Amendment – Change in</li> </ul>	11/23/99	010	Original Protocol	7/10/97
	Protocol M/3342/0004 Amendment 11		012	Amendment 1	8/22/97
			015	Amendment 2 and 3	9/30/97
			018	Amendment 4 and 5	2/4/98
			022	Amendment A, 6 and 7	4/1/98
			025	Amendment 8	9/2/98
			027	Amendment 9	1/30/98
·			039	Amendment 10	12/9/98
090	Protocol Amendment –	12/22/99	Ç,		
	New Investigators and new laboratory to M/3342/0016		049	Original protocol	5/10/99
	<ul> <li>New sub-investigators for Protocol</li> </ul>		041	Original Protocol	2/5/99
	M/3342/0006		043	Amendment 1	3/29/99
			053	Amendment 2	8/20/99
	Information Amendment –				
	Pharmacology/Toxicology Data		N/A	N/A	N/A
150	1 21	12/20/00	000		11/12/07
100	<ul> <li>Frotocol Amenament – Clinical Study</li> </ul>	17/73/39	000	Original Protocol	11/13/96
	Reports		001	Amendment 1	12/12/96
	<ul> <li>a0063183 (Protocol M/3342/0001)</li> </ul>		005	Amendment 2	1/7/97
	<ul> <li>a0062071 (protocol M/3342/0011)</li> </ul>		034	Final Protocol	10/5/98

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
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062	•	Protocol Amendment - Clinical Study	1/18/00	049	Original protocol	5/10/99
		Report and New Investigators to Protocol M/3342/0016		054	Amendment 1	9/21/99
063	•	Annual Report – October 1, 1998 – September 30, 1999	2/15/00	N/A	N/A	N/A
064	•	Protocol Amendment - New	2/18/00	049	Original protocol	5/10/99
		Investigators for M/3342/0016		054	Amendment 1	9/21/99
	•	Information Amendment – Toxicology Reports		N/A	N/A	N/A
990	•	Protocol Amendment - New	3/17/00	049	Original protocol	5/10/99
		Investigators new sub-investigators and new satellite for Protocol M/3342/0016		054	Amendment 1	9/21/99
	•	Information Amendment -		N/A	N/A	N/A
		<ul><li>Clinical Report</li><li>Toxicology Report</li></ul>				
990	•	Protocol Amendment - Change in	3/21/00	041	Original Protocol	2/5/99
		Protocol M/3342/0006 Amendment 3		043	Amendment 1	3/29/99
	$\downarrow$			000	ל זווסוווסווע	0/20/22
190	•	Protocol Amendment - New	4/18/00	049	Original protocol	5/10/99
		Investigators and sub-investigators for		054	Amendment 1	9/21/99
	•	Information Amendment – Toxic reports		N/A	N/A	N/A
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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
890	General Correspondence – Transfer of Sponsorship from Pharmacia & Upjohn Company to Boehringer Ingelheim Pharmaceuticals, Inc.	4/19/00	N/A	N/A	N/A
690	General Correspondence – Boehringer     Ingelheim's acknowledgment of transfer     of IND Sponsorship	4/19/00	890	Pharmacia & Upjohn's General Correspondence – Transfer of Sponsorship	4/19/00
070	<ul> <li>Protocol Amendment – New Investigator for M/3342/0016</li> </ul>	2/30/00	049 054	Original protocol Amendment 1	5/10/99 9/21/99
	Information Amendments – CMC and Pharmacology/Toxicology Data		N/A	N/A	N/A
071	<ul> <li>Protocol Amendment – Change in Protocol M/3342/0004 (BI Trial 1182.1)</li> </ul>	8/3/00	010 012	Original Protocol Amendment 1	7/10/97 8/22/97
			015	Amendment 2 and 3	9/30/97
			018	Amendment 4 and 5	2/4/98
			022	Amendment 6 and 7	4/1/98
			025	Amendment 8	96/2/98
			027	Amendment 9	1/30/98
			039	Amendment 10	12/9/98
			029	Amendment 11	11/23/99
072	Protocol Amend – New Protocol 1182.5	9/13/00	N/A	N/A	N/A

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
No			Submission	Referenced	of Reference	
073	•	Protocol Amendment – New Investigators for:	10/6/01			
		• 1182.5		072	Original Protocol	9/13/00
		• 1182.1 (M/3342/0004)		010	Original Protocol (1182.1)	7/10/97
	•	Updated FDA Form 1572 for Investigator in BI Trial 1182.7 (M/3342/0016)		049 054	Original protocol Amendment 1	5/10/99 9/21/99
074	•	Response to FDA Comments dated	00/9/11	N/A	Fax from FDA	9/13/00
		comments by the Medial Reviewer and Medical Team Leader regarding Protocol		071	Protocol Amendment – Change in Protocol M/3342/0004/BI Trial	8/3/00
		Amendment – Change in Protocol M/3342/0004/BI Trial 1182.1			1182.1	
075	•	Request for Type B Meeting to discuss clinical development plans for tipranavir	12/27/00	N/A	Teleconference with FDA	12/21/00
920	•	Protocol Amendment – New Protocol and New Investigator for 1182.6	10/61/1	N/A	N/A	N/A
077	•	General Correspondence – Background Document for End of Phase I Meeting	2/9/01	075	Request for Type B Meeting	12/27/00
		scheduled for 4/5/01		N/A	DAVDP Advisory Committee Meeting	1/11/01
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078	•	Protocol Amendment -	2/12/01	N/A	N/A	N/A
		<ul> <li>New protocol for 1182.17</li> </ul>		049	Original protocol	66/01/5
		<ul> <li>Change in Protocol 1182.4</li> </ul>		054	Amendment 1	9/21/99
		Amendment 2				
		<ul> <li>Change in Protocol 1182.6 -</li> </ul>		920	Original protocol	1/19/01
		Amendment 1				
020	•	Annual Report – October 1, 1999 – September 30, 2000	2/13/01	N/A	N/A	N/A
080	•	Protocol Amendment - Change in	2/23/01	049	Original protocol	5/10/99
<del></del>		Protocol 1182.6 Amendment 2		054	Amendment 1	9/21/99
081	•	Information Amendment - Clinical Trial	3/7/01	N/A	N/A	N/A
		Keports				
085	•	Response to FDA comments faxed on	3/16/01	N/A	Fax with FDA Comments	3/1 and 5/01
		March $3 - 5$ , 2001 regarding 1182.6			regarding 1182.6	
- u <del>-</del>				9/0	Original Protocol	1/19/01
				7.20	End of Phase I Background	2/9/01
					Document	
083	•	General Correspondence - Update to	3/30/01	710	End of Phase I Background	2/9/01
		Background Document for End of Phase			Document	
	<del></del>	I Meeting scheduled for April 5, 2001		N/A		3/29/01
					Telephone conservation between BIPI and FDA	
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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
S <sub>o</sub>	_		Submission	Referenced	of Reference	
084	•	Response to FDA Comments regarding	4/3/01	072	Fax with FDA Comments	11/9/00
085	•	Response to FDA Comments regarding	4/3/01	078	Fax with FDA Comments	3/19/01
		BI Trial No. 182.17			regarding 1182.17	
980	•	Information Amendment – Clinical Trial	4/4/01	N/A	N/A	N/A
087	Ļ	Demonse to EDA Comments recording	4/4/01	N/N	Telenhone convercation hetween	1/3/01
200	•	BI's upcoming Pre-IND meeting	10/ <del>†</del> / <del>†</del>	V/N	BIPI and FDA regarding	10/6/4
					upcoming pre-IND meeting	
880	•	General Correspondence - Meeting	4/19/01	N/A	End of phase I Meeting	4/5/01
		Minutes from End of Phase I Meeting				
680	•	IND Safety Report –	4/24/01	N/A	N/A	N/A
		<ul> <li>Initial Report 2001-BP-01258</li> </ul>			;	
060	•	Protocol Amendment - New	5/7/01			
		Investigators for:				
	•	1182.2		041	Original Protocol	2/5/99
				043	Amendment 1	3/29/99
				053	Amendment 2	8/20/99
				990	Amendment 4	3/21/00
				I		
	•	1182.6		076	Original Protocol	1/19/01
				078	Amendment 1	2/12/01
				080	Amendment 2	2/23/01
		• 1182.17		078	Original Protocol	2/12/01
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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
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091	•	IND Safety Report	5/10/01	N/A	N/A	N/A
	_	<ul> <li>Initial 2001-BP-01439</li> </ul>				
092	•	IND Safety Report	5/25/01			
		<ul> <li>Initial Report 2001-BP-01639</li> </ul>	•	N/A	N/A	N/A
		• Follow-up #1 2001-BP-01439		091	Initial Report	5/10/01
093	•	IND Safety Report Initial 2001-DE05034	5/29/01	N/A	N/A	N/A
094	•	Protocol Amendment - Changes in	5/31/01			
		protocol				
		<ul> <li>1182.2 Amendment 4</li> </ul>		041	Original Protocol	2/5/99
				043	Amendment 1	3/29/99
				053	Amendment 2	8/20/99
				990	Amendment 3	3/21/00
-	-					
		<ul> <li>1182.4 Amendment 3</li> </ul>		049	Original Protocol	2/10/99
<u>.</u>				054	Amendment 1	9/21/99
				078	Amendment 2	2/12/00
095	<u> •</u>	General Correspondence – requesting	6/6/01	N/A	N/A	N/A
		FDA concurrence with designation of				
		starting materials in the syntheses of				
	4	tipranavir drug substance				
960	•	Protocol Amendment - New				
		Investigators for:				
-,		• 1182.6		920	Original Protocol	1/19/01
				078	Amendment 1	2/12/01

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
No			Submission	Referenced	of Reference	
				080	Amendment 2	2/23/01
		• 1182.17		078	Original Protocol	2/12/01
097	•	IND Safety Report –	6/12/01	160	Initial Report	5/10/01
į		<ul> <li>Follow-up #2 2001-BP-01439</li> </ul>		092	Follow-up #\1	5/25/01
860	•	IND Safety Report –	6/12/01	092	Initial Report	5/25/01
		<ul> <li>Follow-up #1 2001-BP-01639</li> </ul>				
660	•	Request for Type C Meeting to discuss	6/14/01	N/A	End of phase I Meeting with	4/5/01
		cillical developinent plans		088		4/19/01
					Draft Meeting minutes from End	
_					of Phase I Meeting	
				N/A	,	6/8/01
					Telephone conversation between BIPI and FDA	
100	•	Information Amendment – CMC support for on-going and future clinical trails	6/19/01	N/A	N/A	N/A
101	•	Protocol Amendment – Changes in 1182.17 Amendment 1	6/25/01	078	Original Protocol	2/12/01
102	•	IND Safety Reports	7/3/01			
		<ul> <li>Initial Report 2001-FF-C0384</li> </ul>		N/A	N/A	N/A
		<ul> <li>Follow-up Report #3 2001-BP-01439</li> </ul>		091	Initial Report	5/10/01
		•		092	Follow-up #1	5/25/01
				600	Follow-up #2	6/12/01
	<del> </del>	• Follow-up Report #2 2001-BP-01639		092	Initial Report	5/25/01
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103   Description of Submission   Date of Serial No.   Description of Submission   Serial No.   Description of Submission   Stating Reference   of Referen		A.A.	A WALL T	A A PARMITA NOT THE A WILLIAM A SAN AND AND AND AND AND AND AND AND AND A	1001 LVB	
• IND Safety Report Initial Report         7/25/01         N/A         N/A         N/A           • IND Safety Report Initial Report         7/27/01         076         Original Protocol           • Protocol Amendment - New Investigators         7/27/01         076         Original Protocol           • I182.6         Amendment 1         080         Amendment 1           • IND Safety Report         8/1/01         093         Initial Report           • General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance         8/3/01         095         General Correspondence - requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance           • IND Safety Report         N/A         Teleconference between FDA and BI           • IND Safety Report         103         Initial Report	Serial	Description of Submission	Date of	Serial No.	Description of Deference	Date of Reference
• IND Safety Report Initial Report         7/25/01         N/A         N/A           • Protocol Amendment - New Investigators         7/27/01         076         Original Protocol           • 1182.6         • 1182.17         Amendment 1         080         Amendment 2           • IND Safety Report         078         Original Protocol         Amendment 1           • IND Safety Report         8/1/01         093         Initial Report           • Follow-up # 1         2001-DE-05034         8/3/01         095         General Correspondence – the definition of starting materials in synthesis of TPV drug substance           • The definition of starting materials in synthesis of TPV drug substance         N/A         Teleconference between FDA and BI           • IND Safety Report         8/8/01         103         Initial Report           • Follow-up Report #1 2001-DB-00029         103         Initial Report				860	Follow-up #1	6/12/01
• Protocol Amendment - New Investigators         7/27/01         076         Original Protocol           • 1182.6         • 1182.0         Amendment 1         080         Amendment 2           • 1182.17         078         Original Protocol         078         Original Protocol           • IND Safety Report         8/1/01         093         Initial Report           • Follow-up # 1         2001-DE-05034         8/3/01         095         General Correspondence - tequesting FDA concurrence with designation of starting materials in synthesis of TPV drug substance           • General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance         N/A         Teleconference between FDA and BI           • IND Safety Report         BI         Initial Report           • Follow-up Report #1 2001-DB-00029         Initial Report         Initial Report	103	IND Safety Report Initial Report 2001- DB-00029	7/25/01	N/A	N/A	N/A
<ul> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>Follow-up # 1 2001-DE-05034</li> <li>General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance</li> <li>IND Safety Report</li> <li>INA</li> <li>Initial Report</li> <li>General Correspondence - requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance</li> <li>IN/A</li> <li>Initial Report</li> <li>Initial Report</li> </ul>	104	<ul> <li>Protocol Amendment - New Investigators</li> <li>1182.6</li> </ul>	1/27/01	076 078 080	Original Protocol Amendment 1 Amendment 2	1/19/01 2/12/01 2/23/01
<ul> <li>IND Safety Report</li> <li>Follow-up # 1</li> <li>General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance</li> <li>Initial Report</li> <li>General Correspondence - requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance</li> <li>N/A</li> <li>IN/A</li> <li>Teleconference between FDA and BI</li> <li>Initial Report</li> <li>Follow-up Report #1 2001-DB-00029</li> <li>Initial Report</li> </ul>		• 1182.17		078 101	Original Protocol Amendment 1	2/2/01 6/25/01
<ul> <li>General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance</li> <li>IND Safety Report</li> <li>Follow-up Report #1 2001-DB-00029</li> <li>General Correspondence - requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance</li> <li>N/A</li> <li>Teleconference between FDA and BI</li> <li>Initial Report</li> </ul>	105		8/1/01	093	Initial Report	5/29/01
• IND Safety Report • Follow-up Report #1 2001-DB-00029	106	General Correspondence - Proposal for the definition of starting materials in synthesis of TPV drug substance	8/3/01	095 N/A	General Correspondence – requesting FDA concurrence with designation of starting materials in the syntheses of tipranavir drug substance Teleconference between FDA and BI	6/6/01
	107	IND Safety Report     Follow-up Report #1 2001-DB-00029	8/8/01	103	Initial Report	7/25/01

Serial Description of Submission  No  108  • IND Safety Reports  • Follow-Up Report #4 2001-BP-  109  • Request for Type C Meeting - Req FDA feedback on adequacy of our selection approach.  110  • Follow-up Report  • Follow-up Report  • Follow-up Report  • Follow-up Report  111  • Protocol Amendment - Changes in Protocol 1182.6 Amendment 3  112  • Follow-up Report  114  • Follow-up Reports			24		SOF PORTED STATES AND A CONTROL OF A	20 TACE	
•   •   •   •   •   •   •   •   •   •	Seria	lal	Description of Submission	Date of	Serial No.	Description	Date of Reference
	No	_		Submission	Referenced	of Reference	
• • • •	108	∞	IND Safety Reports	8/14/01	:		
• • • •			<ul> <li>Follow-Up Report #4 2001-BP-01439</li> </ul>		160	Initial Report	5/10/01
• • • •			•		092	Follow-up #1	5/25/01
• • • •				_	260	Follow-up #2	6/12/01
• • • •					102	Follow-up #3	7/3/01
• • • •			• Follow-Up Report #3 2001-BP-0639		092	Initial Report	5/25/01
• • • •			T T		860	Follow-up # 1	6/12/01
• • • • •					102	Follow-up # 2	7/3/01
• • • •	109	6	Request for Type C Meeting - Requesting	8/17/01	N/A	End of Phase I Meeting	4/5/01
• • • •			FDA feedback on adequacy of our dose				
• • • •			selection approach.		880	End of Phase I Meeting Minutes	4/19/01
• • •	110	0	IND Safety Report	8/20/01	103	Initial Report	7/25/01
• • •			<ul> <li>Follow-up Report #2 2001-DB-00029</li> </ul>		107	Follow-up #1	8/8/01
• • •	111	1	<ul> <li>Protocol Amendment - Changes in</li> </ul>	8/20/01	920	Original Protocol	1/19/01
• •			Protocol 1182.6 Amendment 3		078	Amendment 1	2/12/01
• • •					080	Amendment 2	2/23/01
• •	112	2	IND Safety Report	8/27/01	102	Initial Report	7/3/01
• •			<ul> <li>Follow-up Report #1 2001-FF-C0384</li> </ul>				
•	113	3	<ul> <li>Protocol Amendment - New Investigators</li> </ul>	8/29/01	078	Original	2/12/01
•			for 1182.17		101	Amendment 1	6/25/01
Follow-up Report #5 2001-BP-	114	4	<ul> <li>IND Safety Reports</li> </ul>	8/30/01	091	Initial Report	5/10/01
			• Follow-up Report #5 2001-BP-01439		092	Follow-up #1	5/25/01
				• • •	260	Follow-up #2	6/12/01
					102	Follow-up #3	7/3/01

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
%		Submission	Referenced	of Reference	
			108	Follow-up #4	8/14/01
115	Desk Copies of General Correspondence     Background Document for Type C     Meeting	9/25/01	109	General Correspondence - Background Document for Type C Meeting	8/17/01
116	Response to FDA Request for Information – Diskette with MS Word Files from Background Document for Type C Meeting	9/26/01	109	General Correspondence - Background Document for Type C Meeting	8/17/01
				General Correspondence - Background Document for Type C Meeting	
117	Response to FDA Request for Information containing a chart with the Overview of Tipranavir Clinical Trial Program with Particular Focus on Subjects 1182.12 and 1182.48	9/27/01	N/A	N/A	N/A
118	IND Safety Reports     Follow-in Report #4 2001-BP-01639	9/28/01	092	Initial Report Follow-up # 1	5/25/01
			102	Follow-up # 2 Follow-up # 3	7/3/01 8/14/01
119	<ul><li>IND Safety Report</li><li>Initial Report 2001-FF-C0647</li></ul>	10/12/01	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
120	Information Amendment -     Pharmacology/Toxicology	10/12/01	N/A	N/A	N/A
121	<ul><li>Protocol Amendment</li><li>New Protocols 1182.37 and 1182.41</li></ul>	10/30/01	N/A	N/A	N/A
	• Changes in Protocol 1182.37		N/A	N/A	N/A
	New Investigators for 1182.6		076 078	Original Protocol Amendment 1	1/19/01 2/12/01
			080	Amendment 2 Amendment 3	2/23/01 8/20/01
-	New Investigators for 1182 17		078	Original	2/12/01
	TOTAL TOTAL PROPERTY OF THE TAXABLE TOTAL TAXABLE TAXA		101	Amendment 1	6/25/01
122	<ul> <li>General Correspondence - Request for Advice on Duration of Combination Toxicology Studies</li> </ul>	11/7/01	N/A	N/A	N/A
123	General Correspondence - Meeting     Minutes from October 5, 2001 meeting     discussing BI's clinical Development     plans and Interaction Program	11/7/01	N/A	FDA and BI Meeting	10/5/01
124	IND Safety Report     Follow-in Report #5 2001-BP-01639	11/8/01	092	Initial Report Follow-up # 1	5/25/01 6/12/01
	COOL TO TOO TO TOO TO TOO TO TOO TO TOO TO	-	102	Follow-up # 2	7/3/01
			108	Follow-up # 3	8/14/01
			118	Follow-up # 4	9/28/01

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Š		Submission	Referenced	of Reference	
125	<ul> <li>Protocol Amendment – Changes in Protocol</li> </ul>	11/13/01			
	• 1182.37 Amendment 2		121	Original Protocol and Amendment	10/30/01
	• 1182.41 Amendment 1		121	Original Protocol	10/30/01
:	<ul> <li>Information Amendment - Clinical Report, Tipranavir Investigators Brochure</li> </ul>				
126	Response to FDA Request for  The formation is greated to formation.	11/20/01	N/A	Fax from FDA	11/16/01
	information in regards to fax requesting sample patient informed consent form		121	Amendment 1 (1182.37)	10/30/01
	and investigators documentation for Protocols 1182.37 and 1182.41		125	Amendment 2 (1182.37)	11/13/01
			121 125	Original Protocol (1182.41) Amendment 1 (1182.41)	10/30/01 11/13/01
127	<ul> <li>Serial Number Correction to Information Amendment – Clinical/Clinical Pharmacokinetics</li> </ul>	12/4/01	N/A	Information Amendment – Clinical/Clinical Pharmacokinetics	11/30/01
128	• IND Annual Report – Reporting period October 1, 2000 – September 28, 2001	12/14/01	N/A	N/A	N/A
129	General Correspondence – Discontinuation of Tipranavir Study 1182.42	12/19/01	N/A	Telephone conversation between BI and FDA	12/18/01

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
130	<ul> <li>Protocol Amendment - Changes in</li> </ul>	12/19/01	121	Original Protocol	10/30/01
	Protocol 1182.37 Amendment 3		121	Amendment 1	10/30/01
			125	Amendment 2	11/13/01
131	<ul> <li>IND Safety Report Follow-up #2</li> </ul>	12/20/01	093	Initial Report	5/29/01
	2001-DE-05034(2)		105	Follow-up #1	8/1/01
132	Information Amendment - Clinical     Pharmacokinetics from 1182 6	12/20/01	N/A	Clinical Development Meeting between BI and FDA	10/2/01
			920		1/19/01
_			820	Original Protocol	2/12/01
			080	Amendment 1	2/23/01
			111	Amendment 2	8/20/01
				Amendment 3	
133	Response to FDA Comments in regards	12/27/01	N/A	FDA Fax with Comments	11/15/01
	to comonation toxicology studies		N/A	Telephone conversations between FDA and BI	12/12-13/01
134	Protocol Amendment -	12/27/01			
	• Changes in Protocol 1182.41		121	Original Protocol	10/30/01
	Amendments 2 and 3		123	Amenument 1	11/13/01
	New Investigators in Protocol		078	Original Protocol	2/12/01
	1182.17		101	Amendment 1	6/25/01

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
No	_		Submission	Referenced	of Reference	
135	•	IND Safety Report	12/28/01	102	Initial Report	7/3/01
		<ul> <li>Follow-up Report #2 2001-FF-C0384</li> </ul>		112	Follow-up Report #1	8/27/01
136	•	Response to FDA Comments in regards	1/29/02	N/A	Telephone conversations between	1/28-1/29/02
		to formulation on tipranavir used in			BI and FDA	
		preclinical toxicology studies		100	Information Amendment - CMC	6/19/01
	•	Information Amendment-			support for on-going and future	
		Pharmacology/Toxicology/Chemistry,			clinical trails studies.	
		Manufacturing and Control	•	133	Response to FDA Comments in	12/27/01
					regards to combination toxicology studies	
137	•	Information Amendment:	2/14/02	129	Terminated study	12/19/01
		Clinical/Request for Comment-				
		Discontinuation of Tipranavir Study				
		1182.42				
138	•	Protocol Amendment	3/4/02	N/A	Clinical Development Meeting	10/5/01
		<ul> <li>New Protocol 1182.12</li> </ul>			between BI and FDA	
		<ul> <li>Changes in Protocol 1182.12</li> </ul>				
		Amendment 1				
139	•	Response to FDA Request for	3/5/02	N/A	FDA Fax	2/15/02
		Information				
140		<ul> <li>Information Amendment -</li> </ul>	3/13/02	N/A	N/A	N/A
		Pharmacology/Toxicology Reports				
141	•	General Corresp – BI seeking	3/21/02	960	General Correspondence –	6/6/01
		concurrence with designation of 2			requesting FDA concurrence with	
		starting materials in the synthesis of TPV			designation of starting materials in	

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
	tipranavir drug substance			the syntheses of tipranavir drug substance	
142	Protocol Amendment	3/26/02			
	<ul> <li>New Protocol 1182.55</li> </ul>		N/A	N/A	N/A
	Changes in Protocol 1182.55     Amendment 1				
	New Investigator for 1182.17		078 101	Original protocol Amendment 1	2/12/01 6/25/01
143	Information Amendment - Chemistry,     Manufacturing and Control reports	4/3/02	N/A	N/A	N/A
144	Information Amendment- Pharmacology/Toxicology Reports	4/11/02	N/A	N/A	N/A
145	IND Safety Report	4/24/02	5	1 - 32 - 1	10,01
	• Follow-up Report #3 2001-FF-C0384(3)		102 112 135	initial Keport Follow-up Report #1 Follow-up Report #2	//3/01 8/27/01 12/28/01
· · · · · · · · · · · · · · · · · · ·	• Follow-up Report #1 2001-FF-C0647(1)	4/24/02	119	Initial Report	410/15/01
146	Information Amendment – Clinical     Investigators Brochure Version 5	4/26/02	N/A	N/A	N/A
147	IND Safety Report Follow-up Report #4 2001-FF-0384(4)	4/29/02	102	Initial Report	7/3/01

Date of Serial No.   Description   Submission   Submission   Submission   Submission   Submission   Referenced   of Reference   of Reference   of Reference   of Reference   of Reference   of Reference   112   Follow-up Report #1   12   Follow-up Report #2   12   12   12   12   145   Follow-up Report #3   4/2   Follow-up Report #3   4/				1	9,	
Submission   Referenced   Of Reference   Submission   Reference   Of Reference	Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
112   Follow-up Report #1   113   Follow-up Report #1   114   Follow-up Report #2   145   Follow-up Report #3   145   Follow-up Report #3   145   Follow-up Report #3   145   Follow-up Report #3   145   Follow-up Report #1	N <sub>0</sub>		Submission	Referenced	of Reference	
• IND Safety Report         135         Follow-up Report #2           • Follow-up Report #3         577/02         093         Initial Report #3           • Follow-up Report #3         105         Follow-up Report #1         131           • Follow-up Report #3         105         Follow-up Report #1           • Follow-up Report #2         119         Initial Report           2001-FF-C0647(2)         145         Follow-up Report#1           • Protocol Amendment - Changes in Protocol I 182.12 Amendment 2         5/15/02         138         Original Protocol Amendment 1           • Response to FDA Request for the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials         5/12/02         048         Request for Carcinogenicity Study Protocol           • Information Amendment: Pharmacology/Toxicology report Pharmaco				112	Follow-up Report #1	8/27/01
• IND Safety Report         5/7/02         Follow-up Report #3         Follow-up Report #3           • Follow-up Report         3/7/02         093         Initial Report           • Follow-up Report #3         131         Follow-up Report#1           • Follow-up Report#2         119         Initial Report           • Protocol Amendment - Changes in Protocol Amendment 2         119         Initial Report           • Protocol Amendment - Changes in Protocol I182.12 Amendment 2         138         Original Protocol           • Response to FDA Request for Information - providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical trials         5/12/102         N/A         Response to FDA Request for Information Amendment: Information Amendment: Animacology/Toxicology report         5/24/02         048         Request for Carcinogenicity Study Protocol           • Information Amendment: Pharmacology/Toxicology report         5/24/02         048         Request for Carcinogenicity Study Protocol           • Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year         N/A         FDA Fax				135	Follow-up Report #2	12/28/01
<ul> <li>IND Safety Report</li> <li>Follow-up Report #3</li> <li>Follow-up Report #3</li> <li>Follow-up Report#1</li> <li>Follow-up Report#1</li> <li>Follow-up Report#1</li> <li>Follow-up Report#1</li> <li>Protocol Amendment - Changes in Protocol I 182.12 Amendment 2</li> <li>Response to FDA Request for Information - providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical trials</li> <li>Information Amendment:</li> <li>Information A</li></ul>				145	Follow-up Report #3	4/24/02
Follow-up Report #3      Follow-up Report #3      Follow-up Report #3      Follow-up Report #1      Follow-up Report	148	IND Safety Report	5/7/02			
• Follow-up Report#2  • Follow-up Report#2  • Potocol Amendment - Changes in Protocol Amendment 2  • Protocol 182.12 Amendment 2  • Response to FDA Request for toxicology report request for the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy 135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  • Information Amendment:  • Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year		<ul> <li>Follow-up Report #3</li> </ul>		093	Initial Report	5/29/01
<ul> <li>Follow-up Report#2  2001-FF-C0647(2)  Protocol Amendment - Changes in Protocol I 182.12 Amendment 2  Response to FDA Request for Information - providing information on going six-month rat and dog toxicology studies, the oral toxicity of polyoxy 135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  Information Amendment:  Informatio</li></ul>		2001-DE-05034(3)	•	105	Follow-up Report#1	8/1/01
Follow-up Report#2      Protocol Amendment - Changes in Protocol Amendment - Changes in Protocol 1182.12 Amendment 2      Response to FDA Request for Information - providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials     Information Amendment:     Request for FDA Feedback - BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year      Request PA Request for FDA Fax      Initial Report     Follow-up Report#1     Follow-up Report#1     Amendment Initial Protocol     Information between FDA and BI Request for Information between Information Pharmacology/Toxicology report     Request for FDA Feedback - BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year    Initial Report Follow-up Report#1     Amendment Information between Information between Information between Information between Information Pharmacology/Toxicology report Information Amendment:    Initial Report   Follow-up Request for PDA Request for PDA Fax			-	131	Follow-up Report#2	12/20/01
• Protocol Amendment - Changes in Protocol 1182.12 Amendment 2  • Protocol Amendment 2  • Protocol Amendment 2  • Response to FDA Request for Information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  • Information Amendment:  • Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year  • Protocol 1182.12 Amendment 1  Amendment 1  Amendment 1  Amendment 1  FDA and BI  Response to FDA Request for Enchanger for Information  Request for Carcinogenicity Study  FDA Fax		Follow in Denorth?		119	Initial Report	10/15/01
<ul> <li>Protocol Amendment - Changes in Protocol Amendment 2 Protocol 1182.12 Amendment 2 Protocol 1182.12 Amendment 2</li> <li>Response to FDA Request for Information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials</li> <li>Information Amendment:  Pharmacology/Toxicology report  Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year</li> </ul>		2001-FF-C0647(2)		145	Follow-up Report#1	4/22/02
<ul> <li>Response to FDA Request for Information – providing information – providing information on going six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials</li> <li>Information Amendment:  Pharmacology/Toxicology report  Request for FDA Request for Carcinogenicity Study Protocol  Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year</li> </ul>	149	Protocol Amendment - Changes in	5/15/02	138	Original Protocol	3/4/02
<ul> <li>Response to FDA Request for Information – providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials</li> <li>Information Amendment: 5/24/02 048 Request for Carcinogenicity Study Protocol</li> <li>Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year</li> </ul>		Protocol 1182.12 Amendment 2			Amendment 1	3/4/02
Information – providing information on the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  Information  SEDDS formulations used for other drugs in preclinical/clinical trials  Information  Information  SA24/02  Information  Request for Carcinogenicity Study Protocol  Request for FDA Feedback – BI seeking FDA Fax  The TPV/RTV and RTV groups for 2-year  The TPV/RTV and RTV groups for 2-year	150	<ul> <li>Response to FDA Request for</li> </ul>	5/17/02	N/A	Telephone conversation between	2/15/02
the in-life data at 3 months from the ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  Information  SEDDS formulations used for other drugs in preclinical/clinical trials  Information  SEDDS formulations used for other drugs in preclinical/clinical trials  Information  SEDDS formulations used for other drugs in preclinical/clinical trials  FRACOCCOL  Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year the TPV/RTV and RTV groups for 2-year		Information - providing information on			FDA and BI	
ongoing six-month rat and dog toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  Information Amendment:  Pharmacology/Toxicology report  Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year		the in-life data at 3 months from the				
toxicology studies, the oral toxicity of polyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  Information Amendment: Pharmacology/Toxicology report Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year		ongoing six-month rat and dog		139	Response to FDA Request for	3/5/02
bolyoxy135 castor oil in animals and SEDDS formulations used for other drugs in preclinical/clinical trials  Information Amendment: Pharmacology/Toxicology report Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year		toxicology studies, the oral toxicity of			Information	
SEDDS formulations used for other  drugs in preclinical/clinical trials  Information Amendment:  Pharmacology/Toxicology report  Request for Carcinogenicity Study Pharmacology/Toxicology report  Request for FDA Feedback – BI seeking FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year		polyoxy135 castor oil in animals and				
<ul> <li>drugs in preclinical/clinical trials</li> <li>Information Amendment:         <ul> <li>Information Amendment:</li></ul></li></ul>		SEDDS formulations used for other				
<ul> <li>Information Amendment:         Pharmacology/Toxicology report         Request for Carcinogenicity Study         Protocol         Request for Carcinogenicity Study         Protocol         Request for Carcinogenicity Study         Protocol         FDA concurrence with dose selection for the TPV/RTV and RTV groups for 2-year     </li> </ul>		drugs in preclinical/clinical trials				
Protocol N/A FDA Fax	151	<ul> <li>Information Amendment:</li> </ul>	5/24/02	048	Request for Carcinogenicity Study	5/19/99
N/A FDA Fax		Pharmacology/Toxicology report			Protocol	
N/A FDA Fax		<ul> <li>Request for FDA Feedback – BI seeking</li> </ul>				
the TPV/RTV and RTV groups for 2-year		FDA concurrence with dose selection for		N/A	FDA Fax	5/19/01
		the TPV/RTV and RTV groups for 2-year				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>o</sub>		Submission	Referenced	of Reference	
	rat carcinogenicity study		N/A	Telephone conversation between BI and FDA	4/18/02
152	Protocol Amendment - New Investigators	5/24/02	138	Original Protocol and Amendment	3/4/02
	• 1182.52		149	1 Amendment 2	5/15/02
	• 1182.55		142		3/26/02
				Original Protocol and Amendment 1	
153	IND Safety Report	5/31/02	660	Initial Report	5/29/01
	Follow-up Report #4		105	Follow-up Report #1	8/1/01
	2001-DE-05034(4)		131	Follow-up Report #2	12/20/01
			148	Follow-up Report #3	5/7/01
154	<ul> <li>Protocol Amendment - Changes in</li> </ul>	6/12/02	820	Original Protocol	2/12/01
	Protocol 1182.17 Amendment 2		101	Amendment 1	6/25/01
155	<ul> <li>Protocol Amendment - New Investigators</li> </ul>	6/14/02	138	Original Protocol and Amendment	3/4/02
	for 1182.52		149	1	5/15/02
				Amendment 2	
156	<ul> <li>Information Amendment - Chemistry,</li> </ul>	6/28/02	N/A	N/A	N/A
	Manufacturing and Control				
157	Protocol Amendment-Changes in     Drotocol 1182 55 Amendment 2 and 3	7/2/02	142	Original Protocol and Amendment	3/26/02
	11000001 102.33 Amelianient Z and 3		i	ı	
158	IND Safety Report     Initial Pancer 2003 BE 00410EE(0)	7/9/02	N/A	N/A	N/A
	IIIIIIIII KEPOIL 2002-FF-00410FF(0)				

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
No	_		Submission	Referenced	of Reference	
159	•	Information Amendment-Chemistry, Manufacturing and Control	7/15/02	N/A	N/A	N/A
160	•	General Correspondence - Discontinuation of Tipranavir Study 1182.22	20/11/1	142 157	1182.55 Original Protocol and Amendment 1 Amendments 2 and 3	3/26/02 7/2/02
				138 149	1182.52 Original Protocol and Amendment 1	3/4/02
				149	Amendment 2	5/15/02
				078 101	1182.55 Original Protocol Amendment 1	2/12/01 6/25/01
				152	Amendment 2	6/12/02
161	•	Request for Special Protocol Assessment- Clinical BI requests special protocol assessment of 1182.12	7/18/02	N/A	BI and FDA meet for Type C Meeting	10/5/01
				115	Desk Copies of General Correspondence - Background Document for Type C Meeting	9/18/01
				138	Original Protocol	3/4/02
				149	Amendment 1 Amendment 2	5/15/02

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Keterenced	of Keterence	
162	Information Amendment-	7/19/02	151	Information Amendment –	5/24/02
	Pharmacology/Toxicology Reports			Pharmacology/Toxicology	
	<ul> <li>Request for FDA</li> </ul>				
	Feedback/Teleconference BI request		N/A	FDA Fax accepting proposed dose	6/25/02
	concurrence with lowering dose level of			levels	
	ritonavir		N/A		6/15/02
				BI email to FDA requesting rapid	
				agreement on ritonavir dose in study	
163	IND Safety Report	7/23/02	158	Initial Report	7/9/02
	<ul> <li>Follow-up Report #1</li> </ul>				
	2002-FF-0041OFF (1)				
164	IND Safety Report	8/1/02	158	Initial Report	7/9/02
. <u>-</u>	<ul> <li>Follow-up Report #2</li> </ul>		163	Follow-up #1	7/23/02
	2002-FF-00410FF (2)				
165	<ul> <li>IND Safety Reports</li> </ul>	8/12/02	158	Initial Report	70/6/1
	<ul> <li>Follow-up Report #3</li> </ul>		163	Follow-up #1	7/23/02
	2002-FF-00410ff(3)		164	Follow-up #2	8/1/02
	• Initial Report 2002-RP-03620RP(0)		A/N	7-Dav fax	8/5/02
166	Protocol Amendment –	8/15/02	142	Original Protocol and Amendment	3/26/02
	New Investigator for 1182 55		157		7/2/02
	Changes in Protocol 1182.55			Amendment 2 and 3	
	Amendment 4				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
2	į	Submission	Referenced	of Reference	
167	<ul> <li>IND Safety Reports</li> </ul>	8/23/02	N/A	N/A	N/A
	<ul> <li>Initial and Follow-up #1</li> </ul>				
	2002-FF-00494FF(0)/(1)				
168	IND Safety Report	8/27/02	167	Initial Report	8/23/02
	• Follow-up report #2 2002-FF-00494FF(2)		167	Follow-up #1	8/23/02
169	IND Safety Report	8/28/02	158	Initial Report	7/9/02
	Follow-up Report #4		163	Follow-up #1	7/23/02
	2002-FF-00410FF(4)		164	Follow-up #2	8/1/02
			165	Follow-up #3	8/12/02
170	IND Safety Report	70/9/6	158	Initial Report	7/9/02
	<ul> <li>Follow-up Report #5</li> </ul>		163	Follow-up#1	7/23/02
	2002-FF-00410FF(5)		164	Follow-up#2	8/1/02
			165	Follow-up#3	8/12/02
			169	Follow-up#4	8/28/02
	• Follow-up Report #1		165	Initial Donout	0/17/0
	2002-BP-03629BP(1)		103	niitiai nepoit	0/17/02
171	IND Safety Report	9/12/02	119	Initial Report	10/15/01
	• Follow-up #3 2001-FF-C0647(3)		145	Follow-up #1	4/24/02
			148	Follow-up #2	5/7/02
172	IND Safety Report	9/18/02	167	Initial Report	8/23/02
	Follow-up #3 2002-FF-00494FF(3)		167	Follow-up #1	8/23/02
			168	Follow-up #2	8/27/02

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S <sub>o</sub>		Submission	Referenced	of Reference	
173	IND Safety Report	9/20/05	165	Initial Report	8/12/02
	<ul> <li>Follow-up Report #2</li> </ul>		170	Follow-up#1	9/6/02
	2002-BP-03629BP(2)				
	<ul> <li>Follow-up Report #3 2002-BP-03629BP(3)</li> </ul>	-			
174	General Correspondence –	9/20/02	141	General Correspondence – BI	3/21/02
	Administrative Information for			seeking concurrence with	
	September 27, 2002 teleconference			designation of two starting	
	between BI and FDA to discuss BI's			materials in the synthesis of	
	proposal for designation of 4 TPV as a			tipranavir drug substance	
	starting material in the synthesis of		N/A		8/2/02
	tipranavir drug substance			BI fax to FDA	
			N/A		8/29/02
				FDA Letter re: comments on	9/20/02
				1182.12 and drug interaction	
				program	
175	<ul> <li>Information Amendment-</li> </ul>	10/8/02	N/A	N/A	N/A
	Pharmacology/Toxicology Reports				
176	<ul> <li>Request for Type B Meeting - End of</li> </ul>	10/11/02	N/A	BI and FDA Meeting -Type C	10/5/01
	Phase II				
	<ul> <li>Request for Teleconference-</li> </ul>		123	Meeting minutes issued	11/7/01
	Clinical/Pharmacokinetics				
			138	Protocol 1182.52 submitted	3/4/02
177	<ul> <li>Request for Type B Meeting - End of Phase II (CMC)</li> </ul>	10/11/02	N/A	Meeting Request	

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S S		Submission	Referenced	of Reference	
178	IND Safety Reports	10/16/02			
	<ul> <li>Initial Report 2002-FF-00587FF(0)</li> </ul>		N/A	N/A	N/A
	• Follow-in #4 2002-FF-00494FF(4)		167	Initial Report	8/23/02
			167	Follow-up #1	8/23/02
			168	Follow-up #2	8/27/02
			172	Follow-up #3	9/18/02
179	IND Safety Report	10/24/02	N/A	N/A	N/A
	<ul> <li>Initial Report 2002-IT-00096IT(0)</li> </ul>			-	
180	IND Safety Report	10/29/02			
	• Follow-up #4 2002-BP-03629BP(4)	-	165	Initial Report	8/12/02
	•		170	Follow-up # 1	20/9/6
			173	Follow-up # 2	9/20/02
			173	Follow-up # 3	9/20/02
				!	9
	<ul> <li>Follow-up #6 2002-FF-00410FF(6)</li> </ul>	-	158	Initial Report	7/9/02
			163	Follow-up # 1	7/23/02
			164	Follow-up # 2	8/1/02
			165	Follow-up # 3	8/12/02
			169	Follow-up # 4	8/28/02
			170	Follow-up # 5	9/6/02
181	<ul> <li>Response to FDA Comments - FDA</li> </ul>	10/30/05	N/A	FDA Letter	8/29/02
	Letters re: Special Protocol Assessment,				
	Protocol 1182.12 and ddI Drug		161	Protocol for Request for Special	7/19/02
	Interaction			Protocol Assessment	

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Serial No	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
			N/A	FDA Fax	9/20/02
			N/A	Telecon between BI and FDA	9/20/02
			137	Information Amendment: Clinical/Request for Comment- Discontinuation of Tipranavir Study 1182.42	10/11/02
			176	Request for Type B Meeting – End of Phase II and Request for teleconference	
182	<ul> <li>General Correspondence – Feedback from IRB's refusing to conduct study</li> <li>Request for Feedback</li> <li>Drug Interaction Study with abacavir</li> </ul>	11/8/02	N/A	FDA and BI Meeting	10/5/01
183	<ul> <li>Information Amendment - Chemistry,</li> <li>Manufacturing and Control</li> </ul>	11/13/02	N/A	N/A	N/A
184	<ul> <li>Type B Background Document for End of Phase II Meeting scheduled for December 18, 2002</li> </ul>	11/14/02	177	Type B End of Phase II meeting request	10/11/02
185	<ul> <li>Information Amendment- Pharmacology/Toxicology Reports</li> </ul>	11/14/02	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
186	General Correspondence - Background     Document for End of Phase II Meeting	11/15/02	176	Request for Type B End of phase 2 Meeting	10/11/02
	. •		N/A	Telephone conversation between BI and FDA	10/31/02
			161	Request for a Special protocol Assessment	7/18/02
			N/A	FDA Fax	8/28/02
			N/A	FDA Fax	9/20/02
			181	Response to FDA Comments - FDA Letters	10/30/02
			N/A	FDA Fax "Overview of Pharmacology/Toxicology comments regarding the clinical development of tipranavir"	10/30/02
187	Response to FDA Comments – Overview of Pharmacology/Toxicology comments	11/18/02	N/A	FDA Fax	10/30/02

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>			Submission	Referenced	of Reference	
		regarding the clinical development program for tipranavir		181	Response to FDA Comments - FDA Letters	10/30/02
188	•	IND Safety Report	11/20/02	167	Initial Report	8/23/02
		<ul> <li>Follow-up #5 2002-FF-00494FF(5)</li> </ul>		167	Follow-up # 1	8/23/02
		•		168	Follow-up # 2	8/27/02
				172	Follow-up # 3	9/18/02
				178	Follow-up # 4	10/16/02
189	•	IND Safety Report	11/20/02	178	Initial Report	10/16/02
		• FOIIOW-UP #1 2002-FF-0058/FF(1)				
190	•	Protocol Amendment-	11/26/02			
		<ul> <li>New Investigators for 1182.17</li> </ul>		078	Original Protocol	2/12/01
				101	Amendment 1	6/25/01
			-	154	Amendment 2	6/12/02
191	•	General Correspondence - Chemistry,	12/11/02	N/A	BI and FDA Teleconference	9/27/02
		providing current specifications for 2		N/A	FDA Meeting Minutes	10/7/02
		TPV and information regarding m-			,	
		nitropropriophenone (m-NPP)		N/A	BI and FDA telephone conversation	7/18/01
192	•	Type B Background Document - Chemistry, Manufacturing and Control	12/11/02	184	Type B Background Document - Chemistry, Manufacturing and	11/14/02
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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
193	IND Safety Report	12/11/02	179	Initial Report	10/24/02
	• Follow-up Report #1 2002-IT-00096IT(1)				
194	General Correspondence - Background     Document for End of Phase 2 meeting	12/13/02	186	General Correspondence - Background Document for End of Phase 2 Meeting	11/15/02
195	Response to FDA Comments – Clinical and Clinical Pharmacology comments	12/13/02	N/A	BI and FDA telephone	12/13/02
	regarding tipranavir				
196	<ul> <li>General Correspondence - Request for FDA Feedback regarding proposal for the</li> </ul>	12/13/02	N/A	BI and FDA teleconference	12/4/02
	study of oral Cremophor EL human exposure				
197	IND Safety Report	12/26/02	165	Initial Report	8/15/02
	• Follow-up #5 2002-BP-0362BP(5)	•	170	Follow-up # 1	9/12/02
			172	Follow-up # 2	9/18/02
			173 180	Follow-up # 3 Follow-up # 4	9/20/02 10/29/02
198	<ul> <li>IND Safety Report-</li> </ul>	12/30/02	N/A	N/A	N/A
	<ul> <li>Initial Report 2002-BP-0617BP(0)</li> </ul>				
199	<ul> <li>Information Amendment-</li> </ul>	12/31/02	N/A	BI and FDA teleconference	12/13/02
	Pharmacology/Toxicology, New				
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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>	_		Submission	Referenced	of Reference	
200	•	IND Annual Report – reporting period October 29, 2001 - October 30, 2002	1/3/03	N/A	N/A	N/A
201	•	IND Safety Report  • Follow-up #1 2002-BP-06174BP(1)	1/10/03	198	Initial Report	12/30/02
202	•	IND Safety Report  • Follow-up #2 2002-BP-06174BP(1)	1/22/03	198	Initial Report Follow-up #1	12/30/02 1/10/03
203	•	General Correspondence –  • Request for Feedback regarding BI's	1/24/03	N/A	End of Phase II Meeting	12/17-18/02
		<ul> <li>approach for the early access program</li> <li>Draft Emergency Use</li> <li>(Compassionate Use) Protocol 1182.XX</li> <li>Draft Expanded Access Protocol</li> <li>Concept Sheet</li> </ul>				
204	•	Protocol Amendment New Protocol 1182.51	1/30/03	N/A	End of Phase II Meeting	12/17/02
205	•	Protocol Amendment – New Protocol 1182.12	2/4/03	186 N/A	End of Phase II Background Document FDA Letter to BI	11/15/02
206	•	General Correspondence- Request for FDA Feedback regarding drug interaction study between tipranavir and atorvastin	2/7/03	N/A	N/A	N/A
207	•	Protocol Amendment – Site Specific Protocol 1182.17 Amendments	2/7/03	190	Protocol Amendment – New Investigators for 1182.17	11/26/02

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
208	<ul> <li>General Correspondence – supply</li> </ul>	2/13/03	N/A	N/A	N/A
	containers having child resistant closures may not be activated				
209	IND Safety Report	2/20/03	167	Initial Report	8/23/02
	<ul> <li>Follow-up #6 2002-FF-00494FF(6)</li> </ul>		167	Follow-up # 1	8/23/02
			168	Follow-up # 2	8/27/02
			172	Follow-up # 3	9/18/02
			178	Follow-up # 4	10/16/02
			188	Follow-up #5	11/10/02
210	<ul> <li>Information Amendment – Clinical</li> </ul>	2/25/03	125	Information Amendment –	11/13/01
	Updated Investigators Brochure Version			Clinical Updated Investigators	
	9			Brochure Version 5	
211	IND Safety Report	3/3/03	198	Initial Report	12/30/02
	• Follow-up #3 2002-BP-06147BP(3)		201	Follow-up #1	1/10/03
			202	Follow-up #2	1/22/03
212	DISCREPENCY WITH SERIAL NO.'S     NO SUBMISSION	N/A	N/A	N/A	N/A
213	General Correspondence – Response to	3/18/03	N/A	FDA Fax	12/31/02
	FDA Fax regarding Clinical, Statistical				
	and Microbiology comments for Protocol			FDA Fax	1/3/03
214	<ul> <li>Protocol Amendment – Changes in</li> </ul>	3/19/03	186	End of Phase II Background	11/15/02
	Protocol 118.12 Amendment 1 and 2			Document	
			205		2/4/03
				Original Protocol	

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
ž	4		Submission	Referenced	of Reference	
215	•	Protocol Amendment – New Investigators for Protocol 1182.12	3/21/03	186	End of Phase II Background Document	11/15/02
		)		205		2/4/03
					Original Protocol	
			-	213		3/19/03
	_				Amendments 1 and 2	
216	•	Request for Special Protocol Assessment	3/24/03	N/A	N/A	N/A
	_	- Carcinogenicity Study Protocol. Bl				
		asking FDA concurrence with the mouse				
217	•	General Correctional process	3/24/03	N/A	End of Dhosa II Meating	12/18/02
i	<u> </u>		CO (F. 7) C	UAI	Lind of 1 hase 11 tweeting	12/10/02
		<ul> <li>BI's End of Phase II Meeting Minutes</li> </ul>				
		<ul> <li>Response to FDA Fax – Request to</li> </ul>				
		change official meeting minutes		N/A	FDA Official Meeting Minutes from End of Phase II Meeting	3/14/03
218	•	IND Safety Reports	3/27/03		8	
	-	Initial Report 2002-BP-01629BP(0)		N/A	A/A	N/A
			•	! ! :		•
		• Follow-up #6 2002-BP-03629BP(0)		165	Initial Report	8/15/02
		•		170	Follow-up # 1	9/12/02
				172	Follow-up # 2	9/18/02
				173	Follow-up # 3	9/20/02
				180	Follow-up # 4	10/29/02
	_			197	Follow-up #5	12/26/02
219	•	IND Safety Report	3/28/03	218	Initial Report	3/27/03
		<ul> <li>Follow-up #1 2003-BP-01629</li> </ul>				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
220	<ul> <li>Protocol Amendment – Changes in Protocol 1182 12 Amendment 3</li> </ul>	3/31/03	186	End of Phase II Background	11/15/02
			205		2/4/02
				Original Protocol	
			214		3/19/02
				Amendment 1 and 2	
			N/A		12/31/02
				Response to FDA comments	
221	<ul> <li>Protocol Amendment – New Protocol</li> </ul>	4/3/03	203	General Correspondence –	1/24/03
	1182.58 Open Label Safety Study			Proposed Emergency Use Program	
222	<ul> <li>Protocol Amendment – Changes in</li> </ul>	4/22/03	820	Original Protocol	2/12/01
	Protocol 1182.17 Amendment 3		101	Amendment 1	6/25/01
			154	Amendment 2	6/12/02
223	<ul> <li>IND Safety Report</li> </ul>	4/24/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-CN-00177CN(0)</li> </ul>				-
224	<ul> <li>IND Safety Report</li> </ul>	4/29/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-BP-02474AU(0)</li> </ul>				
225	<ul> <li>IND Safety Report</li> </ul>	£0/L/S			2
	<ul> <li>Follow-up #1 2003-BP-02474AU(1)</li> </ul>		224	Initial Report	4/29/03
226	<ul> <li>Protocol Amendment – Changes in</li> </ul>	2/6/93	204	Original Protocol	1/30/03
	Protocol 1182.51 Amendment 1 and 2				
227	<ul> <li>Information Amendment – CMC, new</li> </ul>	5/13/04	208	General Correspondence - supply	2/13/03
	documentation for drug substance and			containers having child resistant	
	drug product to support on-going and			closures may not be activated	
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IND Safety Report  Initial Report 2003-DE-01992DE(0)	6/3/03	N/A	N/A	N/A
IND Safety Report  Follow-up #2 2003-BP-02474AU(2)	6/6/03	224 225	Initial Report Follow-up #1	4/29/03 5/7/03
Protocol Amendment - Change in	6/10/03	186	End of Phase II Background	11/15/02
Protocol 1182.12 Amendment 4		306	Document	201710
		607	Original Protocol	70/4/7
		214	•	3/19/02
		N/A	Amendment I and 2	12/31/02
		ć	Response to FDA comments	
		077	Amendment 3	3/31/03
IND Safety Report  Follow-in #1 2003-CN-00177CN(1)	6/13/03	223	Initial Report	4/24/03
Protocol Amendment – New Protocol	6/13/03	N/A	N/A	N/A
Protocol Amendment - Change in Protocol 1182.24	6/18/03	232	Original Protocol	6/18/03
IND Safety Report –  • Follow-up #3 2003-BP-02474AU(3)	6/19/03	224 225 229	Initial Report Follow-up #1 Follow-in #2	4/29/03 5/7/03 6/6/03

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
2		Submission	Referenced	of Reference	
236	IND Safety Report –	6/24/03			
	<ul> <li>Follow-up #4 2003-BP-02474AU(4)</li> </ul>		224	Initial Report	4/29/03
			225	Follow-up #1	5/7/03
			229	Follow-up #2	6/6/03
			235	Follow-up #3	6/19/03
237	<ul> <li>Protocol Amendment – New Investigators for 1182.58</li> </ul>	6/27/03	221	Original Protocol	4/3/03
238	<ul> <li>Protocol Amendment – New</li> </ul>	7/2/03			
	Investigators		186	End of Phase II Background	11/15/02
	• 1182.58		205	Document	2/4/03
		,	214	Original Protocol	3/19/03
		•	220	Amendments 1 and 2	3/31/03
			N/A	Amendment 3	12/31/02
				Response to FDA Comments	
			204		1/30/03
-	• 1182.51		226	Original Protocol	5/9/03
				Amendment 1	
239	<ul> <li>Protocol Amendment – Changes in</li> </ul>	7/9/03	204	Original Protocol	1/30/03
	Protocol 1182.51 Amendment 3		226	Amendment 1 and 2	5/9/03
240	<ul> <li>Response to FDA Request for</li> </ul>	7/14/03	204	Original Protocol	1/30/03
	Information - Comments on Protocol		226	Amendment 1 and 2	5/9/03
	1182.51		239	Amendment 3	7/9/03
		,	N/A	Fax form FDA	4/14/03
241	• IND Safety Report – Initial Report 2003-BP-04595BP(0)	7/22/03	N/A	N/A	N/A

Serial         Description of Submission         Date of Referenced         Pectiform         Description         Date of Referenced         A Referenced         Of Referenced         III.15/02           Pediatric Studies         Proposed Written Agreement for Pediatric Studies         N/A         FDA Pediatric Written Agreement         1/22/03 1/28/03           Request for Teleconference to discuss submission         Request for Teleconference to discuss         N/A         FDA Pediatric Written Agreement         1/21/02           Abmission         Request for Teleconference to discuss submission         N/A         Telephone Conversations between         1/21/02           Abmission         N/A         Telephone Conversations between         1/21/02           Abmission         N/A         Telephone Conversations between         1/21/02           Abmission         Amendment - Telephone Conversations between         1/21/02           Abmission         Amendment - Pediatric Exclusivity Study         8/8/03         242         Amendment 2         4/22/03           Abmitted for Pediatric Exclusivity Study         Request for Teleconference to discuss submission         Request for Teleconference to discuss submission         Request for Teleconference to discuss submission				CONTRACTOR DOTTER TIMETOR		
Submission Referenced of Reference  Proposed Changes in Written Request for 7/23/03 186 General Correspondence – End of Pediatric Studies Proposed Written Agreement for Pediatric Studies Request for Teleconference to discuss Request for Teleconference to discuss Request for Teleconference to discuss IN/A End of Phase II Meeting N/A End	Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Proposed Changes in Written Request for Pediatric Studies     Proposed Changes in Written Request for Proposed Changes in Written Request for Proposed Written Agreement for Pediatric Studies     Request for Teleconference to discuss     N/A End of Phase II Meeting     N/A Telephone Conversations between BI and FDA     Information Amendment – Clinical Protocol Protocol 1182.17, first interim safety analyses     Information Amendment – Pediatric Protocol 8/8/03 078 Original Protocol Amendment 1 154 Amendment 1 154 Amendment 2 222 Amendment 3 222 Amendment 1 Request for Pediatric Studies     Submitted for Pediatric Exclusivity Study Protocol Request for Pediatric Studies	No		Submission	Referenced	of Reference	
Pediatric Studies Proposed Written Agreement for Pediatric Studies N/A Franch Pediatric Written Agreement N/A End of Phase II Meeting N/A Telephone Conversations between Bl and FDA Information Amendment – Clinical Protocol 1182.17, first interim safety analyses Protocol Amendment – Pediatric Protocol Submitted for Pediatric Protocol Submitted for Pediatric Exclusivity Study Submitted for Pediatric Exclusivity Study Intital Report Proposed Written Agreement Request for Pediatric Studies Request for Pedi	242	<ul> <li>Proposed Changes in Written Request for</li> </ul>	7/23/03	186	General Correspondence – End of	11/15/02
Proposed Written Agreement for Pediatric Studies     Request for Teleconference to discuss     Submission     Information Amendment – Clinical Protocol I 182.17, first interim safety analyses     Protocol Amendment – Pediatric Protocol 8/8/03     Protocol Amendment 2     Amendment 3     Amendment 4     Amendment 5     Amendment 5     Amendment 6     Amendment 9		Pediatric Studies			Phase II Background Document,	
Request for Teleconference to discuss     Submission     Information Amendment – Clinical Protocol I 182.17, first interim safety     Protocol Amendment – Pediatric Protocol Submitted for Pediatric Exclusivity Study     InD Safety Report – Information But Submission  Information Amendment – Pediatric Protocol States  Information Amendment – Pediatric Proto		<ul> <li>Proposed Written Agreement for</li> </ul>			Pediatric Proposal	
Request for Teleconference to discuss submission     Information Amendment – Clinical Protocol 1182.17, first interim safety analyses      Protocol Amendment – Pediatric Protocol Submitted for Pediatric Exclusivity Study Submitted for Pediatric Exclusivity Study      IND Safety Report –  Request for Telephone Conversations between BI and FDA  Original Protocol Amendment 1  Amendment 2  Amendment 2  Amendment 3  Proposed Changes in Written Request for Pediatric Studies  Proposed Written Agreement for Pediatric Studies  Request for Teleconference to discuss submission  Initial Report  Initial Report  Initial Report		Pediatric Studies				
Information Amendment – Clinical     Protocol 1182.17, first interim safety     Protocol Amendment – Pediatric Protocol     Submitted for Pediatric Exclusivity Study     Submitted for Pediatric Exclusivity Study     IND Safety Report –     Information Amendment –     Information Amendment – Clinical     Amendment 1     Amendment 2     Amendment 3     Amendment 4     Amendment 5     Amendment 7     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 3     Amendment 3     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 3     Amendment 2     Amendment 3     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 3     Amendment 3     Amendment 3     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 3     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 2     Amendment 3     Amendment 3     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 2     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 2     Amendment 3     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 1     Amendment 1     Amendment 2     Amendment 1     Amendment 1     Amendment 1     Amendment 2     Amendment 2     Amendment 1     Amendment 2     Amendment 3     Amendment 3     Amendment 1     Amendmen		<ul> <li>Request for Teleconference to discuss</li> </ul>		N/A	FDA Pediatric Written Agreement	1/22/03 1/28/03
Information Amendment – Clinical     Protocol 1182.17, first interim safety     Protocol Amendment – Pediatric Protocol     Submitted for Pediatric Exclusivity Study     IND Safety Report –     IND Safety Report –     Initial Report     Initial Report  Information Amendment – Clinical  Initial Report		submission				12/17/02
Information Amendment – Clinical Protocol Protocol 1182.17, first interim safety analyses     Protocol Amendment – Pediatric Protocol Submitted for Pediatric Exclusivity Study Submitted for Pediatric Exclusivity Study      IND Safety Report – Follow-up #1 2003-BP-04595BP(1)      Information Amendment – Pediatric Brotocol Structure				N/A	End of Phase II Meeting	
<ul> <li>Information Amendment – Clinical Protocol Protocol 1182.17, first interim safety analyses</li> <li>Protocol Amendment – Pediatric Protocol 8/8/03 242 Amendment 3</li> <li>Protocol Amendment – Pediatric Protocol 8/8/03 242 Amendment 3</li> <li>Submitted for Pediatric Exclusivity Study Request for Pediatric Studies and Proposed Written Agreement for Pediatric Studies</li> <li>Request for Teleconference to discuss submission</li> <li>Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement for Pediatric Studies</li> <li>Proposed Written Agreement and Proposed Written Agreement and</li></ul>						7/21/03
<ul> <li>Information Amendment – Clinical Protocol Protocol 1182.17, first interim safety analyses</li> <li>Protocol 1182.17, first interim safety analyses</li> <li>Protocol 1182.17, first interim safety analyses</li> <li>Protocol Amendment 1</li></ul>				N/A	Telephone Conversations between	7/23/03
<ul> <li>Information Amendment – Clinical Protocol Protocol 1182.17, first interim safety analyses</li> <li>Protocol 1182.17, first interim safety analyses</li> <li>Protocol Amendment – Pediatric Protocol SW/03 242 Amendment 3</li> <li>Protocol Amendment – Pediatric Exclusivity Study Submitted for Pediatric Exclusivity Study Submitted for Pediatric Exclusivity Study</li> <li>Proposed Changes in Written Request for Pediatric Studies Proposed Written Agreement for Pediatric Studies</li> <li>Request for Teleconference to discuss submission</li> <li>Follow-up #1 2003-BP-04595BP(1)</li> </ul>		The state of the s			BI and FDA	
Protocol 1182.17, first interim safety analyses  Protocol Amendment — Pediatric Protocol Submitted for Pediatric Exclusivity Study Submitted for Pediatric Exclusivity Study  Initial Report  Protocol Amendment — Pediatric Protocol Submitted for Pediatric Exclusivity Study Framedment — Proposed Changes in Written Request for Pediatric Studies Proposed Written Agreement for Pediatric Studies Framedment — Proposed Changes in Written Request for Pediatric Studies Framedment — Proposed Changes in Written Request for Pediatric Studies Framedment — Proposed Changes in Written Request for Pediatric Studies Framedment — Proposed Changes in Written Request for Pediatric Studies Framedment — Proposed Changes in Written Framedment — Proposed Changes in Written Request for Pediatric Studies Framedment — Proposed Changes in Written Framedment — Proposed P	243	<ul> <li>Information Amendment – Clinical</li> </ul>	7/28/03	078	Original Protocol	2/12/01
<ul> <li>analyses</li> <li>Protocol Amendment – Pediatric Protocol</li> <li>Submitted for Pediatric Exclusivity Study</li> <li>Submitted for Pediatric Exclusivity Study</li> <li>Submitted for Pediatric Exclusivity Study</li> <li>Proposed Changes in Written Request for Pediatric Studies</li> <li>Proposed Written Agreement for Pediatric Studies</li> <li>Request for Teleconference to discuss submission</li> <li>Initial Report</li> <li>Follow-up #1 2003-BP-04595BP(1)</li> </ul>		Protocol 1182.17, first interim safety		101	Amendment 1	6/25/01
<ul> <li>Protocol Amendment – Pediatric Protocol         Submitted for Pediatric Exclusivity Study         Submitted for Pediatric Exclusivity Study         Submitted for Pediatric Exclusivity Study         Proposed Changes in Written Request for Pediatric Studies         Proposed Written Agreement for Pediatric Studies         Proposed W</li></ul>		analyses		154	Amendment 2	6/12/02
<ul> <li>Protocol Amendment – Pediatric Protocol</li> <li>Submitted for Pediatric Exclusivity Study</li> <li>Submitted for Pediatric Exclusivity Study</li> <li>Proposed Changes in Written         <ul> <li>Request for Pediatric Studies</li> <li>Proposed Written Agreement for Pediatric Studies</li> <li>Request for Teleconference to discuss submission</li> </ul> </li> <li>IND Safety Report –         <ul> <li>Follow-up #1 2003-BP-04595BP(1)</li> <li>Initial Report</li> </ul> </li> </ul>		The state of the s		222	Amendment 3	4/22/03
Submitted for Pediatric Exclusivity Study  • Proposed Written Agreement for Pediatric Studies  • Proposed Written Agreement for Pediatric Studies  • Request for Teleconference to discuss submission  • IND Safety Report  • Follow-up #1 2003-BP-04595BP(1)	244		8/8/03	242	<ul> <li>Proposed Changes in Written</li> </ul>	7/23/03
IND Safety Report –      IND Safety Report –      Follow-up #1 2003-BP-04595BP(1)      Proposed Written Agreement for Pediatric Studies     Request for Teleconference to discuss submission      Request for Teleconference to discuss submission      Initial Report  Initial Report		Submitted for Pediatric Exclusivity Study			Request for Pediatric Studies	
IND Safety Report — 8/19/03 241 Initial Report      Initial Report    Pediatric Studies					<ul> <li>Proposed Written Agreement</li> </ul>	
Request for Teleconference to discuss submission     IND Safety Report     Follow-up #1 2003-BP-04595BP(1)      Request for Teleconference to discuss submission      Initial Report  Initial Report					for Pediatric Studies	
• IND Safety Report –         8/19/03         241         Initial Report           • Follow-up #1 2003-BP-04595BP(1)         8/19/03         241         Initial Report					Request for Teleconference to	
• IND Safety Report – 8/19/03 241 Initial Report • Follow-up #1 2003-BP-04595BP(1)					discuss submission	
• IND Safety Report – 8/19/03 241 Initial Report • Follow-up #1 2003-BP-04595BP(1)						
• IND Safety Report – 8/19/03 241 Initial Report • Follow-up #1 2003-BP-04595BP(1)						
• Follow-up #1 2003-BP-04595BP(1)	245	IND Safety Report –	8/19/03	241	Initial Report	7/22/03
		<ul> <li>Follow-up #1 2003-BP-04595BP(1)</li> </ul>				

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Serial	_	Description of Submission	Date of	Serial No.	Description	Date of Reference
No	_		Submission	Referenced	of Reference	
246	•	IND Safety Report –	8/29/03	N/A	N/A	N/A
		<ul> <li>Initial Report 2003-BP-06107BP(0)</li> </ul>				
247	•	Protocol Amendment - Changes in	9/4/03	078	Original Protocol	2/12/01
		Protocol 1182.17 Amendment 4		101	Amendment 1	6/25/01
,				154	Amendment 2	6/12/02
				222	Amendment 3	4/22/03
248	•	IND Safety Report	6/6/63			
		<ul> <li>Follow-up Report #2</li> </ul>		241	Initial Report	7/22/03
		2003-BP-04595BP(2)		245	Follow-up #1	8/19/03
249	•	Protocol Amendment - New	9/9/03	186	End of Phase II Background	11/15/02
		Investigators for 1182.58		202	Document	2/4/03
				214	Original Protocol	3/19/03
				220	Amendments 1 and 2	3/31/03
				N/A	Amendment 3	12/31/02
					Response to FDA Comments	
250	•	Protocol Amendment - New	9/12/03			
		Investigators		186	End of Phase II Background	11/15/02
		• 1182.12		205	Document	2/4/02
				214	Original Protocol	3/19/02
				220	Amendment 1 and 2	3/31/03
				N/A	Amendment 3	12/31/02
					Response to FDA comments	
				078		2/12/01
		• 1182.17		101	Original Protocol	6/25/01
				154	Amendment 1	6/12/02
	-			222	Amendment 2	4/22/03

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
		·	247	Amendment 3	9/4/03
				Amendment 4	
			232		6/13/03
				Original Protocol	
	• 1182.24	•	204		1/30/03
			226	Original Protocol	5/9/03
	• 1182.51			Amendment I and 2	
251	IND Safety Report –	9/22/03	N/A	N/A	N/A
	• Initial Report 2003-BP-06679BP9)				
252	<ul> <li>Information Amendment – Clinical BI Trial No. 1182.41</li> </ul>	9/23/03	N/A	N/A	N/A
253	<ul> <li>Protocol Amendment – New Investigator for 1182.24</li> </ul>	9/24/03	250	Telephone between BI and FDA conversation regarding SN 250	9/24/03
	Correction of error in Serial Number			0	
	Assignments		250	Protocol Amendment - New	9/12/03
			251	Investigators	9/22/03
				IND Safety Report Initial Report 2003-BP-06679BP9)	
			252		9/23/03
				Information Amend - Clinical	
254	<ul> <li>Request for Special Protocol Assessment</li> </ul>	9/26/03	N/A	End of Phase II Meeting between	12/17/02
	– Clinical, Naïve Trial 1182.33			BI and FDA	

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>			Submission	Referenced	of Reference	
255	•	Information Amendment –	9/26/03	N/A	BI and FDA Teleconference	12/13/02
		Pharmacology/Toxicology - 26-week				
		Safety Study of TPV/RTV SEDDS in		199	Information Amendment-	12/31/02
		Beagle Dogs – 13 week draft interim			Pharmacology/Toxicology, New	
		report			Protocol	
				196		12/13/03
	············	:			General Correspondence - Request	
					for FDA Feedback regarding	
					proposal for the study of oral Cremophor EL human exposure	
256	•	Information Amendment –New Protocol 1182.45	10/3/03	N/A	N/A	N/A
257	•	IND Safety Report	10/6/03			
		<ul> <li>Initial Report 2003-BP-06917BP(0)</li> </ul>		N/A	N/A	N/A
		• Follow-up #1 2003-BP-06679BP(1)		250	Initial Report	9/22/03
258	•	IND Safety Report	10/16/03	N/A	N/A	N/A
		<ul> <li>Initial Report 2003-BP-07417BP(0)</li> </ul>				
259	•	IND Safety Report	10/21/03	228	Initial Report	6/3/03
		<ul> <li>Follow-up #1 2003-DE-01992DE(1)</li> </ul>				
760	•	Information Amendment - Chemistry,	11/10/03	244	Protocol Amendment - Pediatric	8/11/03
		<ul> <li>Drug product documentation for new</li> </ul>			Exclusivity Study	
		dosage form, oral solution				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
<b>2</b>		Submission	Referenced	of Reference	
261	<ul> <li>IND Safety Reports</li> </ul>	11/10/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-DE-04674DE(0)</li> </ul>				
	<ul> <li>Initial Report 2003-DE-05569DE(0)</li> <li>Initial Report 2003-RP-09346RP(0)</li> </ul>				
262	Protocol Amendment - New	11/13/03	100	Original Drotocal	4/3/03
	Investigators for 1182.58	CO/C1/11	177	Ouginal riotocol	4/3/03
263	Information Amendment – Clinical	11/14/03	N/A	BI and FDA meet for Type C	10/5/01
	BI Trial 1182.52 Clinical Trial Report			Meeting	
					-
			115	Desk Copies of General	9/18/01
				Correspondence - Background Document for Type C Meeting	
			000	e c	60,77
			138	Original Protocol	3/4/02
			149	Amendment 1&Amendment 2	5/15/02
264	IND Safety Report	1/17/03			
	• Follow-up #1 2003-DE-05569DE(1)		261	Initial Report	11/10/03
265	<ul> <li>Information Amendment – Clinical,</li> </ul>	11/20/03	Telecon	FDA tele call requesting a	10/24/03
	Response to FDA Request for			summary of studies on the in vitro	
	information on in vitro Selection of Virus			selection of virus selection of virus	
	Resistant to Tipranavir	•		resist to TPV	
			186	General Correspondence -	11/15/02
				Background Doc for End of Phase	

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
				II Meeting	
566	<ul> <li>IND Safety Report</li> </ul>	11/21/03			
	<ul> <li>Follow-up #1 2003-DE-09236BP(1)</li> </ul>		261	Initial Report	11/10/03
267	IND Safety Report	12/1/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-BP-09788BR(0)</li> </ul>				
268	IND Safety Report	12/3/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-DE-06008GB(0)</li> </ul>				
569	IND Safety Report	12/4/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-BP-09687BP(0)</li> </ul>				
270	IND Safety Report	12/15/03			
	<ul> <li>Follow-up #1 2003-DE-06008GB(1)</li> </ul>	,	268	Initial Report	12/3/03
271	<ul> <li>IND Safety Report</li> </ul>	12/19/03			
	<ul> <li>Follow-up #1 2003-BP-09788BR(1)</li> </ul>		267	Initial Report	12/1/03
272	IND Safety Report	12/23/03	N/A	N/A	N/A
	<ul> <li>Initial Report 2003-BP-10718BP(0)</li> </ul>				
273	<ul> <li>IND Safety Reports</li> </ul>	12/30/03			
	<ul> <li>Follow-up #2 2003-DE-06008GB(2)</li> </ul>		268	Initial Report	12/3/03
			270	Follow-up #1	12/15/03
	יוז מיוי מי מימי נוו מי יון מי		777	Taitio Donout	10/00/00
	• rollow-up #1 2003-BF-10/18BF(1)		. 717	minai nepon	12/23/03
274	IND Safety Report     Initial Bound 2002 DE 00627 DE (0)	12/30/03	N/A	N/A	N/A
	Illinai Nepoli 2003-FF-0002/FF(U)				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
0 N		Submission	Referenced	of Reference	
275	<ul> <li>IND Safety Report</li> </ul>	12/31/03			
	<ul> <li>Follow-up #2 2003-BP-09788BR(2)</li> </ul>		267	Initial Report	12/1/03
			271	Follow-up #1	12/19/03
276	<ul> <li>Information Amendment – Clinical</li> </ul>		161	Special Protocol Assessment for	7/18/02
	Request for FDA Comments on Special			Protocol 1182.12 (RESIST 1)	
	Protocol Assessment for Protocol		141	FDA Comments re: Special	0/00/02
<del></del> .	(1 1010,000) 21,2011			Protocol Assessment for Protocol	701071
				1182.12 (RESIST 1)	
			186	BI Response to FDA Comments	11/15/02
				on Special Protocol Assessment	
				(RESIST 1)	
277	General Correspondence – Safety				
	Information for January 8th telecon		205	Original Protocol (1182.12)	2/4/03
			214	Amendment 1 and 2 (1182.12)	3/19/03
			220	Amendment 3 (1182.12)	3/31/03
			230	Amendment 4 (1182.12)	6/10/03
			078	Original Protocol (1182.17)	2/12/01
			101	Amendment 1 (1182.17)	6/25/01
			154	Amendment 2 (1182.17)	6/12/02
			222	Amendment 3 (1182.17)	4/22/03
			247	Amendment 4 (1182.17)	9/4/03

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Š		Submission	Referenced	of Reference	
278	<ul> <li>General Correspondence – Response to Statistical Comments to Protocol 1182.12</li> </ul>	1/7/04	N/A	FDA Statistical Comments	9/16/03
	(RESIST 2) Amendment 2		205	Original Protocol	2/4/03
			214	Amendment 1 and 2	3/19/03
			220	Amendment 3	3/31/03
			230	Amendment 4	6/10/03
279	<ul> <li>Response to Request for Information – AEs and SAEs</li> </ul>	1/9/04	N/A	Email sent to BI form FDA	1/8/04
			N/A	Teleconference between BI and	1/8/04
			277	T.D.A.	1/7/04
				General Correspondence – Safety Information for January 8 <sup>th</sup> telecon	
280	General Correspondence - Safety	1/12/04	279	Response to Request for	1/9/04
	Information for January 16, 2004  Meeting to discuss adverse event info			Information – AEs and SAEs	
281	<ul><li>IND Safety Report</li><li>Initial Report 2003-BP-10942BP(0)</li></ul>	1/12/04	N/A	N/A	N/A
282	IND Safety Report     Initial 2003-FF-00644FF(0)	1/12/04	N/A	N/A	N/A
283	General Correspondence – Correction of	1/13/04	280	General Correspondence - Safety	1/12/04
	Attachment 1 and 2 of SN 280 and Safety Information for January 16, 2004			Information for January 16, 2004 Meeting to discuss adverse event	
	Meeting			information for tipranavir	

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
ž	_		Submission	Referenced	of Reference	
284	•	Response to FDA Request for Information – Clinical Safety Information	1/14/04	N/A	Telephone conversation between BI and FDA	1/12/04
285	•	Response to FDA Request for Information – Safety Policies and TPV	1/14/04	N/A	Telephone conversation between BI and FDA	1/12/04
		Rick Benefit				
				277	General Correspondence - Safety	1/7/04
					Information for January 8" telecon	
				284	Response to FDA Request for	1/14/04
					Information - Clinical Safety	
	_				Information	
286	•	Protocol Amendment - New	1/19/04	820	Original Protocol	2/12/01
		Investigators for 1182.17		101	Amendment 1	6/25/01
				154	Amendment 2	6/12/02
				222	Amendment 3	4/22/03
				247	Amendment 4	9/4/03
				250	Amendment 5	9/4/03
287	•	Information Amendment - Clinical -	1/19/04	N/A	Teleconference between BI and	12/4/02
		data from human Cremophor EL assay			FDA	
						12/13/02
				196	General Correspondence - Request	
					for FDA Feedback regarding	
					proposal for the study of oral	
					Cremophor EL human exposure	
					Information Among durant	20/20/03
	-				IIIOIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	9/20/03

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
o Z		Submission	Referenced 255	of Reference Pharmacology/Toxicology - 26- week Safety Study of TPV/RTV SEDDS in Beagle Dogs – 13 week draft interim report	
288	<ul><li>IND Safety Report</li><li>Initial Report 2004-BP-00114BP(0)</li></ul>	1/20/04	N/A	7-day facsimile	1/15/04
289	<ul> <li>Clinical Information Amendment – Fatal Cases for Submission</li> </ul>	1/20/04	N/A	Meeting between BI and FDA	1/16/04
	<ul> <li>Initial Report 2003-DE-06498DE(0)</li> <li>Initial Report 2004-BP-00054BP(0)</li> <li>Initial Report 2004-UK-00030UK(0)</li> </ul>		280	General Correspondence – Safety Information for January 16, 2004 Meeting to discuss adverse event information for tipranavir	1/12/04
290	<ul><li>IND Safety Report</li><li>Follow-up #3 2003-DE-06008GB(3)</li></ul>	1/23/04	268 270 273	Initial Follow-up #1 Follow-up #2	12/3/03 12/15/03 12/30/03
291	Protocol Amendment – New Investigators for 1182.58	1/23/04	N/A	BI and FDA meet for Type C Meeting	10/5/01
			115	Desk Copies of General Correspondence - Background Document for Type C Meeting	9/18/01
			138 149	Original Protocol Amendment 1&2	3/4/02 5/15/02

Serial         Description of Submission         Bate of Reference         Submission         Referenced         of Reference         Date of Reference           292         • IND Safety Report         1/26/04         282         Initial Report         1/12/04           293         • IND Safety Report         1/26/04         N/A         N/A         N/A           294         • IND Safety Report         1/27/04         N/A         N/A         N/A           295         • IND Safety Report         1/27/04         N/A         N/A         N/A           296         • IND Safety Report         1/27/04         N/A         N/A         N/A           296         • IND Safety Report         1/30/04         N/A         N/A         N/A           296         • Initial Report 2004-UK-0003-UK(2)         294         Initial Report 2004-UK-0005-UK         N/A           297         • Initial Report 2004-UK-0005-UK         1/30/04         N/A         N/A         N/A           298         • Information Amendment – Clinical         2/2/04         N/A         N/A         N/A           299         • General Correspondence – drafted         2/2/04         N/A         N/A         N/A           299         • General Correspondence – drafted<						
Submission         Referenced         of Reference           • IND Safety Report         1/26/04         282         Initial Report           • IND Safety Report         1/26/04         N/A         N/A           • Initial Report 2004-FF-00026FF(0)         1/26/04         N/A         N/A           • IND Safety Report         1/27/04         N/A         N/A           • IND Safety Report         1/27/04         N/A         N/A           • IND Safety Report         1/27/04         N/A         N/A           • IND Safety Report         1/2003-BP-07128BP(1)         1/27/04         N/A           • Initial Report 2004-UK-00057UK(0)         1/30/04         N/A         N/A           • Initial Report 2004-UK-00057UK(0)         1/30/04         N/A         N/A           • Initial Report 2004-UK-00057UK(0)         1/30/04         N/A         N/A           • Follow-up #1 2003-BP-07128BP(1)         1/30/04         N/A         N/A           • Request for Type B Meeting - Pre-NDA         1/30/04         N/A         N/A           Meeting         22004-UK-00030UK(2)         22/04         N/A         N/A           • 2004-UK-00030UK(2)         22004-UK-00030UK(2)         22/2/04         N/A         N/A           • Contral	Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
<ul> <li>ND Safety Report</li> <li>Follow-up #1 2003-FF-00644FF(1)</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>Initial Report 2004-FF-00026FF(0)</li> <li>IND Safety Report</li> <li>Initial Report 2004-BP-07128BP(0)</li> <li>IND Safety Report</li> <li>Initial Report 2004-BP-07128BP(1)</li> <li>IND Safety Report</li> <li>Initial Report 2004-BP-07128BP(1)</li> <li>IND Safety Report</li> <li>Initial Report 2004-UK-00057UK(0)</li> <li>IND Safety Report</li> <li>Initial Report 2004-UK-00057UK(0)</li> <li>IND Safety Report</li> <li>Initial Report 2004-UK-00057UK(0)</li> <li>Request for Type B Meeting - Pre-NDA</li> <li>I/30/04</li> <l< th=""><th>Ž</th><th></th><th>Submission</th><th>Referenced</th><th>of Reference</th><th></th></l<></ul>	Ž		Submission	Referenced	of Reference	
• Follow-up #1 2003-FF-00644FF(1)         282         Initial Report           • IND Safety Report         1/26/04         N/A         N/A           • Initial Report 2004-FF-00026FF(0)         1/27/04         N/A         N/A           • IND Safety Report         1/27/04         N/A         N/A           • Initial Report 2003-BP-07128BP(0)         1/27/04         N/A         N/A           • Initial Report 2004-BP-00396BP(0)         1/30/04         N/A         N/A           • Initial Report 2004-UK-00057UK(0)         1/30/04         N/A         N/A           • Follow-up #1 2003-BP-07128BP(1)         294         Initial Report           • Follow-up #1 2003-BP-07128BP(1)         294         Initial Report           • Follow-up #1 2003-BP-07128BP(1)         294         Initial Report           • Follow-up #1 2003-BP-07128BP(1)         2/2/04         N/A         N/A           • Request for Type B Meeting - Pre-NDA         1/30/04         N/A         N/A           • Request for Type B Meeting - Pre-NDA         1/30/04         N/A         N/A           • Safety Information         - Cold-UK-00030UK(2)         - 2004-UK-00030UK(2)         - 2004-UK-00030UK(2)           • Cond-UK-00030UK(2)         - Cond-UK-00030UK(2)         - Cond-UK-00030UK(2)         - Cond-UK-00030UK (2) </th <th>292</th> <th><ul> <li>IND Safety Report</li> </ul></th> <th>1/26/04</th> <th></th> <th></th> <th></th>	292	<ul> <li>IND Safety Report</li> </ul>	1/26/04			
<ul> <li>IND Safety Report</li> <li>Intial Report 2004-FF-00026FF(0)</li> <li>IND Safety Report</li> <li>Intial Report 2003-BP-07128BP(0)</li> <li>IND Safety Report</li> <li>Intial Report 2004-BP-00396BP(0)</li> <li>IND Safety Report</li> <li>Intial Report 2004-BP-00396BP(0)</li> <li>IND Safety Report</li> <li>Intial Report 2004-BP-00128BP(1)</li> <li>IND Safety Report</li> <li>Intial Report 2004-BP-00128BP(1)</li> <li>INA</li> <li>INA</li> <li>Intial Report</li> <l< th=""><th></th><td>• Follow-up #1 2003-FF-00644FF(1)</td><td></td><td>282</td><td>Initial Report</td><td>1/12/04</td></l<></ul>		• Follow-up #1 2003-FF-00644FF(1)		282	Initial Report	1/12/04
• Initial Report 2004-FF-00026FF(0)         I/27/04         N/A         N/A           • IND Safety Report         • Initial Report 2003-BP-07128BP(0)         1/27/04         N/A         N/A           • Initial Report 2004-BP-00396BP(0)         1/30/04         N/A         N/A           • Initial Report 2004-UK-00057UK(0)         1/30/04         N/A         N/A           • Initial Report 2004-UK-00057UK(0)         1/30/04         N/A         N/A           • Follow-up #1 2003-BP-07128BP(1)         294         Initial Report           • Follow-up #1 2003-BP-07128BP(1)         1/30/04         N/A         N/A           • Follow-up #1 2003-BP-07128BP(1)         1/30/04         N/A         N/A           • Follow-up #1 2003-UK-2005-UK-20057UK(0)         1/30/04         N/A         N/A           • Request for Type B Meeting - Pre-NDA         1/30/04         N/A         N/A           • Information Amendment - Clinical Safety Information - 2004-UK-20030UK(2)         2/2/04         N/A         BI/FDA Meeting           • Cond-UK-00030UK(2)         2004-UK-20030UK(2)         2/2/04         N/A         BI/FDA Meeting           • General Correspondence - drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting         1/2/04         N/A         1/2/04         N/A         1/2/04	293	IND Safety Report	1/26/04	N/A	N/A	N/A
<ul> <li>IND Safety Report</li> <li>Initial Report 2003-BP-07128BP(0)</li> <li>IND Safety Report</li> <li>INA</li> <li>INA</li></ul>		<ul> <li>Initial Report 2004-FF-00026FF(0)</li> </ul>				
<ul> <li>Intial Report 2003-BP-07128BP(0)</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>Intial Report 2004-BP-00396BP(0)</li> <li>Intial Report 2004-UK-00057UK(0)</li> <li>Follow-up #1 2003-BP-07128BP(1)</li> <li>Request for Type B Meeting - Pre-NDA</li> <li>Information Amendment - Clinical</li> <li>Information</li> <li>Information</li></ul>	294	<ul> <li>IND Safety Report</li> </ul>	1/27/04	N/A	N/A	N/A
<ul> <li>IND Safety Report</li> <li>Initial Report 2004-BP-00396BP(0)</li> <li>IND Safety Report 2004-UK-00057UK(0)</li> <li>Initial Report 2004-UK-00057UK(0)</li> <li>Follow-up #1 2003-BP-07128BP(1)</li> <li>Request for Type B Meeting – Pre-NDA Meeting</li> <li>Information Amendment – Clinical Safety Information</li> <li>Information Amendment – Clinical Safety Report</li> <li>Information Amendment – Clinical Safety Report</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>		<ul> <li>Initial Report 2003-BP-07128BP(0)</li> </ul>				
<ul> <li>Initial Report 2004-BP-00396BP(0)</li> <li>IND Safety Report</li> <li>Initial Report 2004-UK-00057UK(0)</li> <li>Follow-up #1 2003-BP-07128BP(1)</li> <li>Request for Type B Meeting – Pre-NDA</li> <li>Information Amendment – Clinical</li> <li>Information Amendment – Clinical</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>General Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>	295	<ul> <li>IND Safety Report</li> </ul>	1/27/04	N/A	N/A	N/A
<ul> <li>IND Safety Report</li> <li>Initial Report 2004-UK-00057UK(0)</li> <li>Follow-up #1 2003-BP-07128BP(1)</li> <li>Request for Type B Meeting – Pre-NDA Meeting</li> <li>Information Amendment – Clinical Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>3004-UK-00030UK(2)</li> <li>40003-UK-00030UK(2)</li> <li>5004-UK-00030UK(2)</li> <li>6003-UK-00030UK(2)</li> <li>7004-UK-0003UK(2)</li> <li>7</li></ul>		<ul> <li>Initial Report 2004-BP-00396BP(0)</li> </ul>				
<ul> <li>Initial Report 2004-UK-00057UK(0)</li> <li>Follow-up #1 2003-BP-07128BP(1)</li> <li>Request for Type B Meeting - Pre-NDA Meeting</li> <li>Information Amendment - Clinical Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>General Correspondence - drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>InD Safety Report</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>	296	<ul> <li>IND Safety Report</li> </ul>	1/30/04			
• Follow-up #1 2003-BP-07128BP(1)         294         Initial Report           • Request for Type B Meeting - Pre-NDA Meeting         I/30/04         N/A         N/A           • Information Amendment - Clinical Safety Information         2/2/04         N/A         N/A           • 2004-UK-00030UK(2)         2/2/04         N/A         BI/FDA Meeting           • Ceneral Correspondence - drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting         2/2/04         N/A         BI/FDA Meeting           • IND Safety Report         2/2/04         N/A         7 Day Facsimile           • Initial 2003-BP-08944BP(0)         2/2/04         N/A         7 Day Facsimile		<ul> <li>Initial Report 2004-UK-00057UK(0)</li> </ul>			N/A	N/A
<ul> <li>Request for Type B Meeting – Pre-NDA         Meeting         Information Amendment – Clinical         2004-UK-00030UK(2)         Ceneral Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting         InD Safety Report         MIA         N/A         N/A         N/A         BI/FDA Meeting         Procedures (SOPs) regarding adverse event reporting         InD Safety Report         InD Safety Report         Initial 2003-BP-08944BP(0)     </li> </ul>		<ul> <li>Follow-up #1 2003-BP-07128BP(1)</li> </ul>			Initial Report	294
<ul> <li>Meeting</li> <li>Information Amendment – Clinical Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>General Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>IND Safety Report 2/2/04 N/A 7 Day Facsimile</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>	297	<ul> <li>Request for Type B Meeting – Pre-NDA</li> </ul>	1/30/04	N/A	N/A	N/A
<ul> <li>Information Amendment – Clinical Safety Information</li> <li>Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>General Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>		Meeting				
<ul> <li>Information Amendment – Clinical Safety Information</li> <li>Safety Information</li> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>General Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>IND Safety Report</li> <li>IND Safety Report</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>						
<ul> <li>2004-UK-00030UK(2)</li> <li>2004-UK-00030UK(2)</li> <li>General Correspondence – drafted revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>IND Safety Report 2/2/04 N/A 7 Day Facsimile</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>	298	Information Amendment – Clinical Safety Information	2/2/04	N/A	N/A	N/A
<ul> <li>• Construction of the component of the control of the</li></ul>		• 2004-UK-00030UK(2)				
<ul> <li>General Correspondence – drafted 2/2/04 N/A BI/FDA Meeting revision to corporate Standard Operating Procedures (SOPs) regarding adverse event reporting</li> <li>IND Safety Report 2/2/04 N/A 7 Day Facsimile</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>	6	• 2004-UK-00030UK(2)	9			
Procedures (SOPs) regarding adverse event reporting  N/A 7 Day Facsimile  Initial 2003-BP-08944BP(0)	299	General Correspondence – drafted revision to comorate Standard Operating	2/2/04	N/A	BI/FDA Meeting	1/16/04
<ul> <li>event reporting</li> <li>IND Safety Report</li> <li>Initial 2003-BP-08944BP(0)</li> </ul>		Procedures (SOPs) regarding adverse				
• IND Safety Report • Initial 2003-BP-08944BP(0)		event reporting				
• Initial 2003-BP-08944BP(0)	300	<ul> <li>IND Safety Report</li> </ul>	2/2/04	N/A	7 Day Facsimile	1/27/04
		• Initial 2003-BP-08944BP(0)				

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
No			Submission	Referenced	of Reference	
301	•	IND Safety Report	2/5/04	N/A	N/A	N/A
		<ul> <li>Initial 2004-DE-00370GB(0)</li> </ul>				
302	•	General Correspondence – BI Meeting minutes from face to face meeting	2/5/04	N/A	BI/FDA Meeting	1/16/04
303	•	IND Annual Report – Reporting period 10/1/02 – 9/30/03	2/6/04	N/A	N/A	N/A
304	•	IND Safety Report	2/6/04	268	Initial	12/3/03
		• Follow-up #4 2003-DE-06008GB(4)		270	Follow-up #1	12/15/03
				273	Follow-up #2	12/30/03
				290	Follow-up #3	1/23/04
305	•	Information Amendment – Clinical	7/6/04	N/A	BI/FDA Meeting	1/16/04
		Safety Information		200	4	
		• 2003-DE-03576GB(0)	,	784	Response to FDA Request for	1/14/04
		2003-BP-05935BP(0)				
306	•	Protocol Amendment – Changes in Protocol 1182 14 Amendment 1	2/6/04	244	Original Protocol	8/8/03
307	•	General Correspondence – Request for	2/6/04	N/A	N/A	N/A
		Evaluation of Trade name				
308	•	IND Safety Report	2/9/04			
		<ul> <li>Follow-up #1 2003-BP-08944BP(1)</li> </ul>		300	Initial Report	2/2/04
				N/A	7 day facsimile	1/27/04
		• Follow-up #1 2004-BP-00114BP(1)		288	Initial Report	1/20/04
	$\Box$			N/A	7 day facsimile	1/15/04

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
ŝ	_		Submission	Referenced	of Reference	-
309	•	Information Amendment – Clinical Safety Information  • 2003-FF-00246FF(0)	2/9/04	N/A	N/A	N/A
310	•	Information Amendment – Clinical Safety Information  • 2004-BP-0677BP(0)	2/12/04	N/A	N/A	N/
311	•	Information Amendment – Clinical Safety Information January 2004 Monthly Report	2/12/04	N/A	BI/FDA Meeting	N/A
312	•	Information Amendment – Clinical Safety Information; 2003-BP-09677BP(0) • 2003-BP-06681BP(0) • 2004-IT-00012IT(0)/ • 2004-BP-00825BR(0)	2/13/04	284	Response to FDA Request for Information	1/14/04
313	•	IND Safety Report • Follow-up #1 2004-UK-00057UK(1	2/13/04	296	Initial Report	2/13/04
314	•	Request for Type B Meeting CMC Pre- NDA Meeting	2/17/04	260	Information Amendment – CMC Drug product documentation for new dosage form, oral solution	10/29/03
315	•	Information Amendment – Clinical Safety Information • 2004-FF-00088FF(0)	2/18/04	N/A	N/A	N/A

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Serial No	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
	• 2003-BP-09677BP(0)		284 312	Initial Report Follow-up #1	1/14/04 2/13/04
	• 2004-BP-00825BR(0)		312	Initial Report	2/13/04
	• 2004-BP-00677BP(0)		310	Initial Report	2/18/04
316	<ul><li>IND Safety Report</li><li>Follow-up #1 2004-DE-00370GB(1)</li></ul>	2/18/04	301	Initial Report	2/5/04
317	<ul> <li>Information Amendment – Pharm/Tox</li> </ul>	2/20/04	N/A	N/A	N/A
318	<ul> <li>General Correspondence – Request for FDA Feedback in regards to BI's plans for immunotoxicology testing for Tipranavir</li> </ul>	2/20/04	N/A	N/A	N/A
319	Information Amend – Clin Safety Info	2/23/04	N/A	N/A	N/A
320	<ul><li>IND Safety Report</li><li>Follow-up #2 2003-BP-08944BP(2)</li></ul>	2/24/04	308 308	Initial Report Follow-up #1	2/2/04 2/9/04
321	<ul><li>IND Safety Report</li><li>Follow-up #5 2003-DE-06008GB(5)</li></ul>	2/25/04	268 270 273 290 304	Initial Follow-up #1 Follow-up #2 Follow-up #3 Follow-up #4	12/3/03 12/15/03 12/30/03 1/23/04 2/6/04

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
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322	•	Information Amendment – Clinical Safety Information • 2004-IT-00024IT(0)	2/26/04	N/A	N/A	N/A
323	•	IND Safety Report  • Follow-up #1 2003-BP-10942BP(1)	2/27/04	281	Initial Report	1/12/04
324	•	Information Amendment – Pharm/Tox U04-3011	2/27/04	N/A	N/A	N/A
325	•	Information Amendment - Clinical Safety Information	3/1/04	N/A	N/A	N/A
326	•	IND Safety Report Follow-up #2 2004-DE-00370GB(2) Initial 2004-FF-00117FF(0)	3/1/04	300 316	Initial Report Follow-up #1	2/5/04 2/18/04
327	•	IND Safety Report  Initial 2004-SW-00048SW(0)	3/1/04	N/A	7 day facsimile	2/24/04
328	•	Protocol Amendment – New Investigators for 1182.58	3/3/04	221	Original Protocol	4/3/04
329	•	IND Safety Report • Follow-up #1 2004-UK-00057UK(2)	3/5/04	296 313	Initial Report Follow-up #1	1/30/04 2/13/04
	· · - · · · ·	• Follow-up #2 2003-BP-10718BP(2)		272 274	Initial Report Follow-up #1	12/23/03 12/30/03
330	•	Information Amendment - Clinical	3/5/04			

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	Safety Information		N/A	7-day facsimile	2/2/04
	<ul> <li>2004-BP-00677BP9(0)</li> </ul>		310	Initial Report	2/12/04
			315	Follow-up #1	2/18/04
			325	Follow-up #2	3/1/04
			284	Initial Renort	1/14/04
	• 2003-BP-10111BP(0)		1	mina report	1/11/10
331	<ul> <li>Information Amendment – Clinical Safety Information</li> </ul>	3/8/04	N/A	N/A	N/A
332	IND Safety Report	3/9/04	NA	N/A	N/A
	<ul> <li>Initial 2003-BP-068455P(0)</li> </ul>		569	Initial Report	12/4/03
	<ul> <li>Follow-up #1 2003-BP-09687BP(1)</li> </ul>				
333	IND Safety Report	3/9/04	N/A	N/A	N/A
	<ul> <li>Initial 2003-BP-01512AU(0)</li> </ul>		261	Initial Report	11/10/03
	• Follow-up #1 2003-DE-04674DE(1)				
334	Information Amendment – Clinical Safety Information	3/10/04	N/A	N/A	N/A
335	IND Safety Report	3/10/04	272	Initial Report	12/23/03
	• Follow-up #3 2003-BP-10718BP(3)		274 329	Follow-up #1 Follow-up #2	12/30/03 3/5/04
336	Information Amendment – Clinical	3/12/04	284	Initial Report	1/14/04
	Safety Information		N/A	7-day facsimile	2/23/04
	• 2003-BP-10241BP(1)		325	Initial Report	3/1/04
	• 2003-FF-00105FF(0)				

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
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337	•	Information Amendment – Clinical Safety Information February 2004	3/12/04	N/A	BI/FDA Meeting	1/16/04
		Monthly Report				
338	•	IND Safety Report	3/12/04	300	Initial Report	2/5/04
		<ul> <li>Follow-up #3 2004-DE-00370GB(3)</li> </ul>		316	Follow-up #1	2/18/04
	$\downarrow$			326	Follow-up #2	3/1/04
339	•	Protocol Amendment - New	3/12/04	078	Original Protocol	2/12/01
-		Investigators for 1182.17		101	Amendment 1	6/25/01
				154	Amendment 2	6/12/02
				222	Amendment 3	4/22/03
				247	Amendment 4	9/4/03
340	•	Response to FDA Request – Review of Mortality Rates	3/12/04	N/A	BI/FDA Teleconference	2/19/04
	•	Request for Teleconference to discuss				
		and receive feedback				
;	•					
341	•	IND Safety Report • Follow-up #3 2004-UK-00057UK(3)	3/15/04	296 313	Initial Report Follow-up #1	1/30/04 2/13/04
				329	Follow-up #2	3/5/04
342	•	Information Amendment – Clinical	3/15/04	N/A	N/A	N/A
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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
NO N		Submission	Referenced	of Reference	
343	<ul> <li>Information Amendment – CMC</li> </ul>	3/15/04	N/A	N/A	N/A
	providing new and updated CMC documentation for drug substance				
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344	<ul> <li>Information Amendment – Clinical Safety Information</li> <li>2004-FF-00058FF(0)</li> </ul>	3/16/04	N/A	N/A	N/A
345	Information Amendment – Clinical	3/17/04	N/A	7-day facsimile	3/1/04
	Safety Information	-	331	Initial Report	3/8/04
	• 2004-BP-01447BP(0)		N/A	7-day facsimile	3/15/04
			342	Initial Report	3/15/04
	• 2003-BP-10562BP(0)				
346	General Correspondence – Pre-NDA	3/17/04	297	Request for Pre-NDA Meeting	1/30/04
	Meeting Package		N/A	BI/FDA Telephone conversation	2/12/04
	<ul> <li>Request for Teleconference to discuss</li> </ul>				
	key clinical, statistical and format topics				
347	IND Safety Report	3/18/04	N/A	N/A	N/A
	<ul> <li>Initial 2004-FF-00125FF(0)</li> </ul>				
348	IND Safety Report	3/19/04	332	Initial Report	3/9/04
	• Follow-up #1 2003-BP-06845BP(1)			•	
349	<ul> <li>Information Package for Type B Pre-</li> </ul>	3/19/04	314	Request for Type B Meeting Pre-	2/17/04
	NDA Meeting – CMC			NDA Meeting	
			N/A	CMC End of Phase II Meeting	12/18/02

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Ž		Submission	Referenced	of Reference	
350	Protocol Amendment – Changes in     December 1102 69 Amendment	3/19/04	221	Original Protocol	4/3/03
	1 IOUCOI 1 102.30 AIIICIIUIICII Z		WAI	DI/LDA Meetilig	1/10/
351	Information Amendment – Clinical Safety Information	3/19/04	N/A	N/A	N/A
253	NID C. S.t. D.	7/10/04	2.4.7	1. F. 7. 1. 1. O	707017
325	<ul> <li>IND Safety Report</li> <li>Initial 2004-FF-00125FF(0)</li> </ul>	3/19/04	347	Originally submitted without case narrative and the like analysis report	3/18/04
353	<ul> <li>Information Amendment – Clinical Safety Information</li> </ul>	3/22/04	N/A	N/A	N/A
354	IND Safety Report	3/22/04	268	Initial	12/3/03
	• Follow-up #6 2003-DE-06008GB(6)		270	Follow-up #1	12/15/03
			273	Follow-up #2	12/30/03
			290	Follow-up #3	1/23/04
			304	Follow-up #4	2/6/04
			321	Follow-up #5	2/25/04
355	<ul> <li>Information Amendment – Pharm/Tox</li> </ul>	3/23/04	N/A	N/A	N/A
	U04-3013, U04-3034-02, U04-3035, U04-3036, U04-3037				
356	Information Amendment – Clinical	3/23/04			
	Safety Information				
	• 2004-BP-01804BP(0)		N/A	N/A	N/A
	2004-BP-00953BR(0)		N/A	7 day fax	3/16/04

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Serial		Description of Submission	Date of	Serial No.	Description	Date of Reference
2 Z			Submission	Referenced	of Reference	
357	• In:	Information Amendment – Clinical U04-3025, U04-3026 and U04-3027	3/23/04	N/A	N/A	N/A
358	• In	Information Amendment – Clinical Safety Information	3/24/04	N/A	N/A	N/A
359	- Z	IND Safety Report	3/24/04			
	•	<ul> <li>Follow-up #2 2004-BP-00114BP(2)</li> </ul>		288 308	Initial Report Follow-up #1	1/20/04 2/9/04
·	•	Follow-up #3 2003-BP-04595BP(3)		241	Initial Report	7/22/03
				245	Follow-up #1	8/19/03
			<del>.</del>	248	Follow-up #2	9/9/03
360	- E	IND Safety Report	3/26/04			
	•	• Follow-up #2 2003-BP-10942BP(2)		281	Initial Report	1/12/04
				,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Follow-up #1	2/27/04
				323		
361	• Sa	Information Amendment – Clinical Safety Information • 2003-BP-06941AU(1)	3/29/04	N/A	7-day fax	3/3/04
				342	Initial Report	3/15/04
	•	2004-FF-00058FF(0)	, , , ,	N/A	7-day fax	3/10/04
				344	Initial Report	3/16/04
				353	Follow-up #1 – ck on this	3/22/04
362	Z •	IND Safety Report	3/30/04	347	Initial Report	3/18/04
	•	<ul> <li>Follow-up #1 2004-FF-00125FF(1)</li> </ul>		352	Replacement of Initial Report	3/19/04

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
0Z		Submission	Referenced	of Reference	
363	IND Safety Report	3/31/04	257	Initial Report	10/6/03
	• Follow-up #1 2003-BP-06917BP(1)				
364	IND Safety Report	3/31/04	333	Initial Report	3/9/04
	<ul> <li>Follow-up #1 2003-P-01512AU(1)</li> </ul>				
365	<ul> <li>Response to FDA Request for</li> </ul>	4/1/04	N/A	Telephone conversation	3/17/04
	Information - Follow-up to Mortality		340	Response to FDA Request –	3/12/04
	Analysis – CD4 Cell counts/causes of deaths			Review of Mortality Rates	
366	Information Amendment – Clinical	4/1/04	N/A	N/A	N/A
	Safety Information				
367	Information Amendment – Clinical	4/6/04	N/A	N/A	N/A
	Safety Information				
368	IND Safety Report	4/6/04	241	Initial Report	7/22/03
	<ul> <li>Follow-up #4 2003-BP-04595(4)</li> </ul>		245	Follow-up #1	8/19/03
			248	Follow-up #2	6/6/63
			359	Follow-up #3	3/24/04
369	General Correspondence -	4/6/04	N/A	BI/FDA Meeting	1/16/04
	Background/Current Safety Reporting		302	BI Meeting Minutes	2/5/04
	Processes and Procedures and Proposed		N/A	FDA Meeting Minutes	3/3/04
	updates to Safety Reporting Processes		285	Overview of BI's Safety SOP	1/14/04
	and Procedures				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
	Request for Teleconference to assure that     BI can implement the new Processes and				
	procedures				
370	Information Amendment – Clinical	4/1/04			
	Safety Information				
	• 2003-FF-00518FF(3)		284	Initial Report	1/14/04
			319	Follow-up #1	2/3/04
	-		370	Follow-up #2	4/7/04
	• 2003-CN-00461CN(1)		284	Initial Report	1/14/04
371	Information Amendment – Clinical Safety Information	4/1/04			
	• 2003-FF-00518FF(3)		284	Initial Report	1/14/04
			319	Follow-up #1	2/3/04
			370	Follow-up #2	4/7/04
	• 2003-CN-00461CN(1)		284	Initial Report	1/14/04
372	IND Safety Report	4/9/04	300	Initial Report	2/2/04
	• 2003-BP-08944BP(3)		308	Follow-up #1	2/9/04
	The state of the s		320	Follow-up #2	2/24/04
373	Information Amendment – Clinical	4/9/04	N/A	N/A	N/A
	Safety Information				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S <sub>o</sub>		Submission	Referenced	of Reference	
374	<ul> <li>Information Amendment – Clinical Safety Information</li> </ul>	4/12/04			
	• 2004-BP-02534BP(0)		N/A	7-day facsimile	4/5/04
	• 2003-FF-00625FF(2)		N/A	7-day facsimile	3/9/04
			342 367	Initial Report Follow-up #1	3/15/04
				•	4/6/04
375	IND Safety Report	4/12/04	N/A	N/A	N/A
	• Initial Report 2003-BP-03027BP(0)				
376	IND Safety Report	4/12/04			
	• Follow-up #1 2003-BP-06107BP(1)		246	Initial Report	8/29/03
377	IND Safety Report	4/12/04	N/A	N/A	N/A
	<ul> <li>Initial Report 2004-BL-00067BL(0)</li> </ul>				
378	Information Amendment – Clinical	4/13/04	N/A	BI/FDA Meeting	1/16/04
	Safety Information March 2004 Monthly Report	-			
379	<ul> <li>Information Amendment – Clinical Safety Information</li> </ul>	4/13/04	N/A	N/A	N/A
380	IND Safety Reports	4/13/04			
	• Follow-up #2 2003-CN-00177CN(2)		223 231	Initial Report Follow-up #1	4/24/03 6/13/03
381	<ul> <li>Amendment to Information Package for Type B Pre-NDA Meeting</li> </ul>	4/13/04	349	Information Package for Type B Pre-NDA Meeting – CMC	3/19/04

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
382	IND Safety Report	4/14/04			
	• Follow-up #3 2004-BP-00114BP(3)		N/A	7-day facsimile	1/15/04
			288	Initial Report	1/20/04
			208	Follow-up #1	2/9/04
			359	Follow-up #2	3/24/04
383	<ul> <li>Information Amendment:Clinical</li> </ul>	4/15/04	383		
	<ul> <li>Interim Study results of pharmacokinetic</li> </ul>	-			
	study 1182.51				
384	Response to Request for Information/Request	4/15/04	N/A	Telefax from FDA	4/8/04
	for Teleconference – Requesting additional				
	information pertaining to the preliminary		384	Response to FDA Fax of	4/15/04
	results from the dual-boosted protease			4/8/04:Preliminary Results from	
	inhibitor pharmacokinetic study, 1182.51			pharmacokinetic study 1182.51	
385	IND Safety Report	4/16/04	333	Initial Report	3/9/04
	Follow-up #2 2003-BP-01512AU(2)		364	Follow-up #1	3/31/04
386	<ul> <li>Information Amendment: Clinical</li> </ul>	4/16/04	210	Information Amendment - Clinical	2/25/03
	<ul> <li>Investigators Brochure - Version 7</li> </ul>			Updated Investigators Brochure	
			125	Version 6	11/13/01
				Information Amendment –	
	•			Clinical Updated Investigators	
				Brochure Version 5	
387	<ul> <li>IND Safety Report Initial Report 2004-FF-00234FF(0)</li> </ul>	4/16/04	N/A	N/A	N/A

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
388	• Information Amendment: Clinical Fatal Cases 2003-FF-00640FF(2)	4/20/04	N/A		
389	IND Safety Report     Initial Report 2004-BP-02706BP(0)	4/19/04	N/A	N/A	N/A
390	Amendment: Clinical Fatal Cases 2003- FF-00640FF(2)	4/20/04	N/A	N/A	N/A
391	• IND Safety Report Follow-up #1 2004-SW-00048SW(1)	4/20/04	327	Initial Report	3/1/04
392	<ul> <li>Response to Request for Information: Tabular Listing of Deaths</li> </ul>	4/22/04		Telefax from FDA	4/8/04
			384	Response to FDA Fax of 4/8/04:Preliminary Results from pharmacokinetic study 1182.51	April 15, 2004
393	<ul> <li>IND Safety Report Initial Report 2004-FF-00256FF(0)</li> </ul>	4/26/04	N/A	N/A	N/A
394	<ul> <li>Information Amendment: Clinical Fatal Cases 2003-FF-00367FF(1), 2003-FF- 00440FF(2), 2003-CN-00318CN(1) and 2003-BP-10111BP(3)</li> </ul>	4/26/04	4/26/04	N/A	N/A
395	IND Safety Report     Initial Report 2004-BP-02978BR9(0)	4/27/04	N/A	N/A	N/A
396	Information Amendment: Clinical Report 1182.22 /U03-3408	4/27/04	N/A	N/A	N/A

On of Submission         Date of Submission         Serial No.           Submission         Referenced 4/28/04         389           2-02706BP(1)         4/28/04         N/A           ment: Clinical Fatal 2004-         4/29/04         N/A           33-BP-10111BP(4), 2004-         280           42004-BP-03100MX(0)         280           accordence: Clinical-Safety         4/30/04         N/A           rendment: Clinical-Safety         4/30/04         N/A           rendment: Clinical-Safety         4/30/04         384           rence: Evaluation of gency regarding TPV         4/30/04         384           am         5/03/04         267           coort         5/03/04         271           coort         275           dment:New Investigators         5/03/04         N/A				<b>Q</b>	D	
Submission   Referenced	Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
IND Safety Report	N <sub>0</sub>		Submission	Referenced	of Reference	
Information Amendment: Clinical Fatal 2004- FF-00176FF(2), 2003-BP-10111BP(4), 2004- BP-02413AU(1) and 2004-BP-03100MIX(0)  Information Amendment: Clinical-Safety Information-April 2004 Quarterly Safety Summary  General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  Information-April 2004 Quarterly Safety Summary  General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  Follow-up 2003-BP-09788BR(3)  Protocol Amendment:New Investigators	397	IND Safety Report Follow-up 2004-BP-02706BP(1)	4/28/04	389	Initial Report	4/19/04
Information Amendment: Clinical-Safety     Information Amendment: Clinical-Safety     Summary     Summary  General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program      IND Safety Report     Follow-up 2003-BP-09788BR(3)     Protocol Amendment:New Investigators     S/03/04     N/A	398	Information Amendment: Clinical Fatal 2004- FF-00176FF(2), 2003-BP-10111BP(4), 2004-	4/29/04	N/A	Meeting between BI and FDA	1/16/04
Information Amendment: Clinical-Safety     Information-April 2004 Quarterly Safety     Summary     Summary  General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program      IND Safety Report     Follow-up 2003-BP-09788BR(3)     Follow-up 2003-BP-09788BR(3)     Protocol Amendment:New Investigators     S/03/04     N/A     N/A		BP-02413AU(1) and 2004-BP-03100MX(0)		280	General Correspondence - Safety	1/12/04
Information Amendment: Clinical-Safety     Information-April 2004 Quarterly Safety     Summary  General Correspondence: Evaluation of     Interactions w/ the agency regarding TPV     Development program      IND Safety Report     Follow-up 2003-BP-09788BR(3)     Follow-up 2003-BP-09788BR(3)     Protocol Amendment:New Investigators     S/03/04     N/A  N/A		•			Information for January 16, 2004	
Information Amendment: Clinical-Safety     Information-April 2004 Quarterly Safety     Summary  General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program      IND Safety Report     Follow-up 2003-BP-09788BR(3)     Follow-up 2003-BP-0978BR(3)     Follow-up 2003-BP-0978BR(3)					Meeting to discuss adverse event information for tipranavir	
Information-April 2004 Quarterly Safety Summary General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3)  • Protocol Amendment:New Investigators 5/03/04  277  8 Protocol Amendment:New Investigators 7/03/04  8 N/A	399	Information Amendment: Clinical-Safety	4/30/04	N/A	Meeting w/FDA where BIPI	1/16/04
Summary  General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 271 275 • Protocol Amendment:New Investigators Summary 384 384 384 567 577 771 772 773 774 774 775 775 775		Information-April 2004 Quarterly Safety			agreed to submit for a period not	
General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 271 272 Protocol Amendment:New Investigators S/03/04 N/A		Summary			less than 1 yr: all deaths for	
General Correspondence: Evaluation of Interactions w/ the agency regarding TPV  Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 271  • Protocol Amendment:New Investigators 5/03/04 N/A					unrelated deaths as CIA cases,	
General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 271 275 • Protocol Amendment:New Investigators S/03/04 N/A					Grade 3 & 4 serious cases on a	
General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 277  • Protocol Amendment:New Investigators S/03/04 N/A					monthly basis, and a safety	
General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 277  • Protocol Amendment:New Investigators S/03/04 N/A					summary on a quarterly basis.	
General Correspondence: Evaluation of Interactions w/ the agency regarding TPV  Development program  • IND Safety Report Follow-up 2003-BP-09788BR(3) 271  • Protocol Amendment:New Investigators 5/03/04 N/A					(beginning on 1/19/04)	
Development program	400	General Correspondence: Evaluation of Interactions w/ the agency regarding TPV	4/30/04		Telefax from FDA	April 8, 2004
• IND Safety Report Follow-up 2003-BP-09788BR(3) 271 275 • Protocol Amendment:New Investigators 5/03/04 N/A		Development program		384	Response to FDA Fax of	April 15, 2004
• IND Safety Report Follow-up 2003-BP-09788BR(3) 271 275 • Protocol Amendment:New Investigators 5/03/04 N/A					4/8/04:Preliminary Results from	
• IND Safety Report Follow-up 2003-BP-09788BR(3) 271 273 • Protocol Amendment:New Investigators 5/03/04 267 275					pharmacokinetic study 1182.51	
Follow-up 2003-BP-09788BR(3) 271  • Protocol Amendment:New Investigators 5/03/04 N/A	401	IND Safety Report	5/03/04	267	Initial report	12/1/03
• Protocol Amendment: New Investigators 5/03/04 N/A		Follow-up 2003-BP-09788BR(3)		271	Follow-up #1	12/19/04
Protocol Amendment: New Investigators 5/03/04 N/A				275	Follow-up #2	12/31/04
	402	<ul> <li>Protocol Amendment:New Investigators</li> </ul>	5/03/04	N/A	BI and FDA meet for Type C	10/5/01
		Study 1182.58			Meeting	

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
0N		Submission	Keterenced	of Reference	
			115	Desk Copies of General Correspondence - Background	9/18/01
		-		Document for Type C Meeting	
			138	Original Protocol	3/4/02
403	Information Amendment: Clinical Fatal	5/4/04	403	N/A	N/A
	Cases 2004-ED-00091ED(U)				
404	<ul> <li>Protocol Amendment – Changes in</li> </ul>	5/5/04	840	Original Protocol	2/12/01
	Protocol 1182.17 Amendment 5		101	Amendment 1	6/25/01
			154	Amendment 2	6/12/02
			222	Amendment 3	4/22/03
			247	Amendment 4	9/4/03
405	Information Amendment:Clinical	5/6/04			
	1182.55	5/6/04	142	Original Protocol	3/26/02
		7/2/02	142	Amendment 1	3/26/02
			158	Amendment 2	3/26/02
			158	Amendment 3	3/26/02
406	General Correspondence: CMC Pre-NDA Meeting Minutes	5/7/04	406	Type B pre-NDA Meeting for tipranavir capsules	4/19/04
407	IND Safety Report	5/10/04			
	Follow-up 2004-FF-00117FF(1)		326	Initial Report	3/1/04

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
2		Submission	Referenced	of Reference	
408	• Information Amendment:Clinical Fatal Cases 2004-BP-01767BP(01)	5/10/04		Follow-up Report CIA Case	
409	• IND Safety Report Follow-up#1 2003-BP-10569BP(1)	5/12/04	284	Initial Report	1/14/04
410	IND Safety Report Initial Report 2004-BP-02041BP(0) Initial Report 2004-BP-02715RA(0) Initial Report 2004-BP-03359BP(0)	5/12/04	N/A	N/A	N/A
411	<ul> <li>General Correspondence - Update to Pre- NDA Meeting Package</li> </ul>	5/13/04	297	Request for Pre-NDA Meeting	1/30/04
			346	General Correspondence – Pre- NDA Meeting Package	3/17/04
412	Information Amendment:Clinical Clinical Safety Information April Monthly report	5/14/04	N/A	Meeting w/FDA where BIPI agreed to submit for a period not less than 1 yr: all deaths for unrelated deaths as CIA cases, Grade 3 & 4 serious cases on a monthly basis, and a safety summary on a quarterly basis. (beginning on 1/19/04)	1/16/04

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Serial No	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
413	IND Safety Report Initial Report 2004-BP-03507BP(0)	5/14/04	N/A	N/A	N/A
414	General Correspondence Update to Analysis Proposal	5/14/04	N/A	Telefax from FDA	4/8/04
	•	4/15/04	384	Response to FDA Fax of 4/8/04:Preliminary Results from pharmacokinetic study 1182.51	
		4/22/04	392	Response to Request for Information: Tabular Listing of Deaths	
		4/30/04	400		
				General Correspondence: Evaluation of Interactions w/ the	
				agency regarding 1 P V Development program	
415	IND Safety Report: Initial Report2004-BP-03519BP(0) Initial Report 2004-BP-01283BR(0)	5/17/04	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
416	2004-BP-03100MX(1)	Submission	N/A	Meeting between BI and FDA	1/16/04
			280	General Correspondence – Safety Information for January 16, 2004 Meeting to discuss adverse event information for tipranavir	1/12/04
417	IND Safety Report: Initial Report 2004-BP-01006BP(0)	5/18/04	N/A	N/A	N/A
418	Clinical Safety Information 2003-BP-10341AU(0)	5/18/04	N/A	Meeting between BI and FDA	1/16/04
	2004-FF-00058FF(3)			General Corr. – Safety Information for January 16, 2004 Mtg to discuss adverse event information for TPV	1/12/04
419	IND Safety Report: Initial Report 2004-FF-00264FF(0) Follow-up 2004-BP-03519BP(1)	5/19/04	N/A 415	N/A Initial report	N/A 5/17/04
420	Clinical Safety Information 2004-BP-02646BP(0) 2004-BP-02646BP(1) 2003-BP-1034AU(1)	5/19/04	N/A	Meeting between BI and FDA General Correspondence – Safety Information for January 16, 2004 Meeting to discuss adverse event information for tipranavir	1/16/04

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
421	IND Safety Report	5/20/04			
	2004-DE-02587DE(0)		N/A	N/A	N/A
	2004-FF-00296FF(0)		N/A	N/A	N/A
422	IND Safety Report:	5/20/04			
	Initial: 2004-FF-00284FF(0) Initial:		N/A	N/A	N/A
	2004-FF-00297FF(0)		N/A	N/A	N/A
	Initial: 2003-DE-06173GB(0)		N/A	N/A	N/A
423	IND Safety Report	5/20/04	N/A	Meeting between BI and FDA	1/16/04
	Clinical Safety Information				
	2003-FF-00508FF(2)			General Correspondence - Safety	1/12/04
				Information for January 16, 2004	
				Meeting to discuss adverse event	
				information for tipranavir	
424	IND Safety Report:	5/20/04	N/A	N/A	
	Initial Report: 2003-FF-00503FF(0)				
425	General Correspondence	5/21/04	346	Pre-NDA Meeting Package	3/17/2004
	Update to pre-NDA meeting				
	CTD-integrated analysis	N/A	N/A	Pre-NDA Meeting Teleconference	May 10, 2004
426	Information Amendment: Clinical Fatal Case	5/21/04	N/A	CIA Cases	
	Initial Report: 2004-BP-03659BP(0)				
	Follow-up #1: 2004-BP-02543BP(1)				
	Follow-up #4: 2004-UK-00030UK(4)				
427	General Correspondence - Request for	5/21/04	369	General Correspondence -	April 6, 2004
	Teleconferenc-Update of Safety Reporting		N/A	Teleconference with FDA	April 8, 2004

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
8		Submission	Referenced	of Reference	
	Procedures				
428					
	Follow-up Report: 2004-BP-01006BP(1)	5/21/04	417	Initial Report	5/18/04
	Initial Report 2004BP-03620BP(0) Initial Report 2003-FF-00595FF(0)				
429	IND Safety Report				
	Initial Report: 2004-BP-03727BP(0)	5/21/04	N/A	N/A	N/A
430	IND Safety Report	5/24/04			
	Follow-up: 2004-BP-03659BP(1)		426	Initial	5/21/04
431	2004-BP-03721BP(0)	5/24/04	N/A	N/A	N/A
432	IND Safety Report:	5/24/04	N/A	N/A	N/A
	2004-BP-03640BP(0)				
	2004-BP-03614BR(0)				
433	IND Safety Report: 2004-BP-03805BP(0)	5/26/04	N/A	N/A	N/A
434	IND Safety Report	5/26/04	413	Initial	5/14/04
	2004-BP-03507BP(1)				
	2004-FF-00296FF(1)		421	Initial	5/20/04
435	IND Safety Report	5/26/04	410	Initial	5/12/04
	2004-BP-03359BP(1)				
436	General Correspondence:	5/26/2004	N/A	Teleconference with Division	3/17/04
	Top Line Phase 3 RESIST Data				5/25/04
437	General Correspondence:	5/26/2004	203	Draft Protocol	1/23/03
	Synopsis of Proposed Expanded Access		221	Final Protocol 1182.58	4/3/03
	Program - BIPI Trial 1182.70		350	Amendment 1	3/19/04
			436	GC: Top Line Resist 1 and 2	4/27/04

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
438	Information Amendment: Clinical Fatal Cases 2004-FF-00180FF(1)	5/27/04	353	Initial	3/22/04
439	2004-BP-03519BP(2)	5/27/04	415	Initial	5/17/04
			419	Follow-up 1	5/19/04
	2004-BP-02706BP(2)		389	Initial	4/19/04
			397	Follow-up 1	4/28/04
	2004-BP-01283BR(1)		415	Initial	5/17/04
440	2004-BP-03727BP(1)	5/27/04	429	Initial	5/21/04
441	General Correspondence: Statistical Analysis	5/27/04	346	Pre-NDA Meeting Package	3/17/04
_	Plan – submitting the summary of the RESIST protocol and amendments.			)	
442	IND Safety Report				
	2004-BP-02414BP(0)	5/28/04	N/A	N/A	N/A
	2004-BP-03928BP(0)		N/A	N/A	N/A
	2004-BP-03862BP(0)		N/A	N/A	N/A
443	Information Amendment: Clinical U04-1256 (1182.11)	5/28/04	N/A	N/A	N/A
444	Information Amendment: Clinical Fatal Cases 2004-BP-03860BP(0)	5/28/04	N/A	7 Day Fax	5/24/04
445	Information Amendment Clinical Food Effect	5/28/04	N/A	request from T. Sinha requesting info on food effects on bioavail of	5/27/04
				SEDDS formul. in humans	
446	Information Amendment: Clinical Bilirubin Lab Results	5/28/04	N/a	e-mail request from T. Sinha – requesting updated bilirubin data	5/27/04

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
				for previous submissions	
			375	<ul> <li>IND Safety Report</li> </ul>	4/12/04
			401	Initial Report 2003-BP- 03027BP(0)	
					5/3/04
				• IND Safety Report Follow-up 2003-BP-09788BR(3)	
447	IND Safety Report	5/28/04	N/A	N/A	
	Initial 2004-FF-00327FF(0)				
	Initial 2004-FF-00315FF(0) Initial 2004-BP-03876AU(0)				
448	Information Amendment: CMC Request for	5/28/04	N/A	N/A	
	Type A Meeting				
449	Information Amendment: Clinical Fatal Case 2004-DE-00489GB(0)	6/1/04	N/A	N/A	
450	IND Safety Reports Initial Reports	6/2/04	N/A	N/A	
	2004-FF-00321FF(0) 2004-FF-00326FF(0)				
451	Information Amendment: Clinical	6/3/04	N/A	N/A	
	U03-3131-01 (1182.6)				
	U04-3100 (1182.10)				
	004-3210 (1102.21).003-3003-01 (1102.24)				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
452	IND Safety Report:	6/3/04	424	Initial	5/20/04
	2004-FF-00284FF(1)		422 422		
	2004-FF-00297FF(1)				
453	Protocol Amendment: New Investigators	6/4/04	N/A	N/A	
	1182.58 Trial				
454	IND Safety Report	40/4/9	293	Initial	1/26/04
	2004-FT00020FF(1)				
455	Information Amendment: Clinical	6/1/04		Initial	
	2004-INL-00039INL				
456	IND Safety Report: Follow-up				
	2003-FF-00595FF(1)	6/1/04	428	Initial	5/21/04
	2003-FF-00640FF(3)		379	Follow-up 1	4-13-04
			390	Follow-up 2	4/20/04
457	IND Safety Report:	6/7/04	N/A	N/A	N/A
	Initial 2004-BP-04048BP(0)			-	
	Initial 2004-BP-04071BP(0)				
	Initial 2004-SW-00154DB(0)				
458	IND Safety Report:	6/8/04	N/A	N/A	N/A
	Initial 2004-BP-04135BP(0)				
	Initial 2004-BP-02885BP(0)				
459	IND Safety Report:	6/9/04			
	Follow-up 2003-DE-6173GB(1)		422	Initial	5/20/04
	Follow-up 2003-BP-09788BR(4)		267	Initial	12/1/03
			271	Follow-up #1	12/19/03
			275	Follow-up #2	12/31/03

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
			401	Follow-up #3	5/3/04
	Follow-up 2004-DE-02587DE(1)		421	Initial	5/20/04
	Follow-up 2003-FF-00640FF(4)		284	Initial	1/14/04
			379	Follow-up #1	4/13/04
			390	Follow-up #2	4/20/04
			456	Follow-up #3	6/1/04
460	Information Amendment: CMC Type A IND	6/11/04			
	Meeting Information Package for Oral Solution		448	A Type A "critical path" IND Meeting request	5/28/04
461	IND Safety Report:				,
	Follow-up 2004BP-01006BP(2)	6/14/04	417	Initial	5/18/04
			428	Follow-up 1	5/21/04
462	Information Amendment: Clinical	6/14/04	N/A	Meeting w/FDA where BIPI	1/16/04
	Safety Information			agreed to submit for a period not	
	May I Monthly report			less than 1 yr: all deaths for	
				unrelated deaths as CIA cases,	
				Grade 3 & 4 serious cases on a	
				monthly basis, and a safety	
				summary on a quarterly basis.	
				(beginning on 1/19/04)	
463	IND Safety Report	6/15/04	N/A	7 Day Ear	70/01/9
403	IND Safety Report Initial 2004-UK-00526UK(0)	9/12/04	N/A	/ Day Fax	6/10/04

Serial No 464	Description of Submission				
	Description of Submission	Date of	Serial No.	Description	Date of Reference
		Submission	Referenced	of Reference	
	IND Safety Report	6/15/04	429	Initial	5/21/04
	Follow-up 2004-BP-03727BP(2)		440	Follow-up #1	5/27/04
465 II	IND Safety Report 2004-FF-00342FF(0)	6/15/04	N/A	N/A	N/A
466 II	IND Safety Report	6/16/04	267	Initial	12/1/03
	Follow-up 2003-BP-09788BR(5)		271	Follow-up #1	12/19/03
			275	Follow-up #2	12/31/03
-			401	Follow-up #3	5/3/04
			459	Follow-up #4	6/9/04
467 L	Information Amendment: Clinical	6/16/04	426	Initial Report	5/21/04
<u>ν</u>	Safety 2004-BP-03659BP(2)		430	Follow-up #1	5/24/04
468 II	IND Safety Report	6/17/04	N/A	N/A	N/A
	Initial 2003-FF-00532FF(0)				
1 469 I	Information Response to FDA Request for Information – clarification regarding safety	6/17/04	N/A	Teleconference between BI and Division	6/1/04
re	report case 2004-BP-02646BP(1) from trial 1182.12				
470 I	Information Amendment:PharmTox	6/17/04	151	Draft Protocol Submitted	5/24/02
2	24- Month oral Rat Carcinogenicity Study				
471 C	General Correspondence – Request for FDA Feedback – Request for Telecon –	6/18/04	346 N/A	Electronic submission proposal	3/17/04
A	Amendment to NDA Electronic Submission			Pre-NDA telecon and meeting	5/10/04 & 6/2/04

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
				Informal e-mail from J. Liang	6/14/04
472	IND Safety Report: Follow-up 2004-BP-03876AU(1)	6/18/04	447	Initial	5/28/04
473	IND Safety Report Initial 2004-BP-04375BP(0)	6/18/04	N/A	N/A	N/A
474	Information Amendment: Pharmacology/Toxicology U04-3111	6/21/04			
	Request for FDA Feedback		318	GC: Request for FDA Feedback on Immunotox Proposal FDA Response FAX	2/20/04
475	IND Safety Report Follow-up 2004-BP-03614BR(1)	6/21/04	432	Initial	5/24/04
476	IND Safety Report Initial 2004-BP-04465BP(0)	6/21/04	N/A	N/A	N/A
477	Amendment to Information Package for Type A Meeting: CMC	6/21/04	448	Type A "critical path" IND Meeting request	May 28, 2004
				Information Amendment: CMC Type A IND Meeting Information Package for Oral Solution	June 11, 2004

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478	IND Safety Report Initial 2004-BP-04500RA(0)	6/21/04	N/A	N/A	N/A
479	IND Safety Report Initial 2004-FF-00364FF(0)	6/22/04	N/A	N/A	N/A
	F0110W-up 2004-BF-03802BF(1)		7447	initial	5/28/04
480	IND Safety Report: Follow-in 2004-BP-02715RA(1)	6/22/04	410	[eilin]	2/12/04
481	IND Safety Report	6/22/04	N/A	N/A	N/A
	Initial 2004-FF-00355FF(0) Initial 2004-BP-04612BP(0)		V/V	<b>A</b> /Z	<b>4</b> /2
482	IND Safety Report	6/22/04	N/A	N/A	N/A
	Initial 2004-SW-00154DB(0)				
483	Type A IND Meeting-Response to FDA Pre-	6/22/04		Facsimile correspondence from	5/22/04
_	Meeting Request for Information-CMC-			Ms. Tanima Sinha	
·	Tipranavir Oral Solution		260	Information Amendment	10/29/03
483a	IND Safety Report	6/23/04	457	Initial	40/L/9
	Follow-up 2004-SW-00154DB(1)				
484	IND Safety Report Initial 2004-BP-04608BP(0)	6/23/04	N/A	N/A	N/A
485	IND Safety Report	6/23/04			
	Follow-up 2004-ES-00091ES(1)		403	Initial	5/4/04
	Follow-up 2004-FF-00315FF(1)		447	Initial	5/28/04
	Follow-up 2004-FF-00326FF(1)		450	Initial	6/2/04
-	Follow-up 2003-FF-00595FF(2)		428	Initial	5/21/04
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02		Submission	Keterenced	of Reference	
486	IND Safety Report Initial 2004-FF-00361FF(0)	6/24/04	N/A	N/A	N/A
487	IND Safety Report Initial 2004-BP-04794BP(0)	6/25/04	N/A	N/A	N/A
488	Information Amendment: Pharmacology/Toxicology U04-3184	6/25/04	N/A	N/A	
489	IND Safety Report Initial 2003-FF-00569FF(0)	6/25/04	N/A	N/A	N/A
490	IND Safety Report Follow-up 2004-FF-00321FF(1)	6/25/04	450	Initial	6/2/04
491	IND Safety Report Follow-up 2004-FF-00117FF(2)	6/25/04	326 407	Initial Follow-up #1	3/1/04
492	IND Safety Report Initial 2004-BP-04724BP(0) Initial 2004-FF-00372FF(0) Follow-up 2003-FF-00508FF(3)	6/25/04	N/A N/A 284 373 423	N/A N/A Initial Follow-up#1 Follow-up#2	N/A N/A 1/14/04 4/9/04 5/20/04
493	IND Safety Report Initial 2004-FF-00371FF(0) Initial 2004-FF-00373FF(0)	6/28/2004	NA	NA	NA
494	IND Safety Report Initial 2004-BP-04770BR(0)	6/28/2004	N/A	N/A	N/A

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7	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
	IND Safety Report Initial Report: 2004-BP-04688BR(0)	6/28/2004	NA	NA	NA
	Information Amendment: Clinical Step 1 of Drug Interaction Analysis (SN 414) Request for Teleconference	6/29/04	384	Response to FDA Request for Information	4/15/04
	4		392	Response to Request for Information: Tabular Listing of Deaths	4/22/04
			400	General Correspondence: Evaluation of Interactions w/ the agency regarding TPV Development program	4/30/04
			414	General Correspondence:Update to Analysis Proposal	5/14/04
-	-		N/A	Telefax from FDA	6/8/2004

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Serial No	Description of Submission	Date of	Serial No. Referenced	Description of Reference	Date of Reference
497	IND safety report Follow-up 2004-FF-00355FF(1)	6/29/2004	481	Initial	6/22/04
498	Protocol Amendment Change in Protocol Amendment 5 and 6 1182.12	6/30/2004	205 214 220 230	New Protocol 1182.12 Amendments 1 &2 Amendment 3 Amendment 4	2/4/2002 3/19/2002 3/31/2002 6/10/2003
499	General Correspondence; BIPI Meeting Minutes of June 7, 2004 – safety reporting processes and procedures.	6/30/04	N/A	June 7, 2004 Meeting	N/A
200	IND Safety Report Initial 2004-CN-00238CN(0) Initial 2004-IT-00093IT(0)	6/30/04	N/A	N/A	N/A
501	General Correspondence/Request for FDA Feedback/Request for Teleconference NDA Clinical Summary Cross Referencing Plan, Presentation of Patient Disposition,	7/1/04	N/A N/A	Pre NDA Meeting with FDA Pre-NDA Meeting with FDA Telegonforance with Mc Sinho	May 10, 2004  June 2, 2004
	Chinear Cuminary megration rean.		521	FDA Project Manager Pre-NDA Meeting Minutes	Juic 28, 2007  July 19, 2004
502	IND Safety Report Follow-up 2004-UK-00526UK(1)	7/1/04	463	Initial	6/15/04

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
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503	IND Safety Report Initial 2004-FF-00149FF(0)	7/1/04	N/A	N//A	N/A
504	IND Safety Report Initial 2004-UK-00597UK(0)	7/2/04	7/2/04	N/A	N/A
505	IND Safety Report	7/2/04	293	Initial	1/26/04
	Follow-up 2004-FF-00026FF(2)		454	Follow-up #1	6/4/04
	Follow-up 2004-FF-00180FF(2)		353	Initial	3/22/04
			438	Follow-up #1	5/27/04
	Initial 2004-BP-04910BP(0)	·	N/A	N/A	N/A
909	IND Safety Report	7/2/04	N/A	N/A	2004-UK-
	Initial: 2004-FF-00401FF(0)				00320UK(1) is
	Initial: 2004-UK-00320UK(1)				initial. Incorrectly
	Initial: 2004-BP-04967BP(0)				labeled follow-up 1
207	IND Safety Report	7/6/04	289	Initial	1/20/04
	Follow-up 2004-UK-00030UK(5)		289	Follow-up #1	1/20/04
			298	Follow-up #2	2/2/04
			298	Follow-up #3	2/2/04
			426	Follow-up #4	5/21/04
	Follow-up 2003-FF-00503FF(2)		424	Initial	5/20/04
			452	Follow-up #1	6/3/04
208	General Correspondence Proposed Expanded	7/7/04	437	General Correspondence	5/26/04
	Access Program BIPI Trial 1182.70			draft protocol synopsis	
209	Protocol Amendment	7/7/04	078	Protocol 1182.17	2/12/2001
	New Investigator 1182.17		113	PA Dr. Coleen Tutton	8/29/2001

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S N		Submission	Referenced	of Reference	
510	IND Safety Report	40/6/L			
	Follow-up 2004-BP-04688BR(1)		495	Initial	6/28/04
	Initial 2004-BP-05113MX(0)		N/A	N/A	N/A
511	IND Safety Report	7/12/04			
	Follow-up 2004-03862BP(2)		442	Initial	5/28/04
			479	Follow-up #1	6/22/04
	Follow-up 2004-BP-04724BP(1)		492	Initial	6/25/04
	Follow-up 2004-FF-00342FF(1)		465	Initial	6/15/04
512	Protocol Amendment	7/12/04	221	Protocol 1182.58 original	4/3/03
	New Investigator 1182.58			submission	
	Response to FDA Request for Information	7/12/04	N/A	Telefax from FDA, Anthony	6/3/04
513	Response to Telefax from FDA, Anthony			ElHage – Requesting info on	
	ElHage - Requesting info on investigators,			investigators and sites for the	
	sites, discont. and reasons discont. for all of			pivotal sites for TPV.	
	the pivotal sites for TPV Trials (1182.12,				
	1182.48 and 1182.14)				
514	U04-3257 Toxicokinetic of Tipranavir in a	7/12/04	N/A		
	13-week oral toxicity study in rats.				
515	IND Safety Report	7/13/04			
	Follow-up: 2003-BP-09788BR(6)		267	Initial	12/01/03
			271	Follow-up #1	12/19/03
			275	Follow-up #2	12/31/03
			401	Follow-up #3	5/03/04
			459	Follow-up #4	6/09/04
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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
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			466	Follow-up #5	6/16/04
	Follow-up: 2004-BP-04612BP(1)		481	Initial	6/22/04
	Initial: 2004-BP-05320BP(0)		N/A	N/A	N/A
	Follow-up: 2004-FF-00026FF(3)		293	Initial	1/26/04
			454	Follow-up #1	6/04/04
			505	Follow-up #2	7/02/04
516	Information Amendment: Clinical (U03-3605-01 & U04-3198)	7/13/04	N/A	N/A	N/A
517	Information Amendment: Clinical-Safety	7/14/04	N/A	Meeting w/FDA where BIPI	N/A
	Information-Jun 2004 Monthly Report			agreed to submit for a period not	
				less than I yr: Grade 3 & 4 serious	
				cases on a monthly basis.	
				(beginning on 1/19/04)	
518	IND Safety Report	7/14/04			
	Initial: 2004-DE-03659DE(0)		N/A	N/A	N/A
	Initial: 2004-FF-00410FF(0)	16	N/A	N/A	N/A
	Follow-up:2004-FF-00373FF(1)		493	Initial	6/28/04
	Follow-up: 2004-UK-00526UK(2)		463	Initial	6/15/04
			502	Follow-up #1	7/01/04
	Follow-up: 2004-BP-02715RA(2)		410	Initial	5/12/04
			480	Follow-up #1	6/22/04
519	IND Safety Report	7/15/04	N/A	N/A	N/A
···	Initial: 2004-BP-05348BR(0) Initial: 2004-BP-05301R A(0)				

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
520	IND Safety Report 2004-CN-00253CN(0)	7/16/04	N/A	N/A	N/A
521	General Correspondence: Pre-NDA Meeting Minutes	7/19/04	N/A	Meeting with FDA (Pre-NDA)	6/2/04
522	General Correspondence: Request for FDA Feedback Point of discussion in telecon was the difference between "end-of-text" tables (Section 15 of reports) and appendices (Appendix 16.1.9.2, referred to as STATDOCS). Submission of examples of these tables and appendices to illustrate the content of the displays and their supportive statistical documentation	7/20/04	N/A	teleconference	7/12/04
523	General Correspondence: Step 1 of Drug Interaction Analysis In preparation for our teleconference with Division t July 21, 2004, a clear tabular representation of the drugs proposed for study in Step 2 of our analysis plan	7/20/04	496	submission provided a descriptive analysis of concomitant medications used in HIV-1 patients who have died during tipranavir studies or have had clinical progression events in the two ongoing RESIST trials ( <i>Step I of the Drug Interaction Analysis Plan</i>	6/29/04
524	IND Safety Report Follow-up: 2004-UK-00597UK(1) Initial: 2004-BL-00130BL(0)	7/20/04	504 N/A	Initial N/A	7/02/04 N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
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525	IND Safety Report	7/21/04			
	2004-FF-00327FF(		447	Initial	5/28/04
	Follow-up: 2004-BP-04724BP(2)		492	Initial	6/25/04
			511	Follow-up #1	7/12/04
526	Information Amendment Clinical Reports (U04-3248, U04-3259)	7/22/04	N/A	N/A	N/A
527	Information Amendment:	7/22/04	N/A	N/A	N/A
	Pharmacology/Toxicology (U03-3565, U03-3153)				
528	IND Safety Report	7/22/04			
	Follow-up: 2004-FF-00410FF(1)		518	Initial	7/14/04
	Follow-up: 2004-FF-00256FF(1)		393	Initial	4/26/04
529	IND Safety Report	7/23/04	N/A	N/A	N/A
	Initial: 2004-DE-03872DE(0) Initial: 2004-BP-05611BP(0)				
530	IND Safety Report:	7/26/04	N/A	N/A	N/A
	Initial: 2004-CN-00263CN(0)				
531	IND Safety Report	7/27/04			
	Follow-up: 2004-DE-03659DE(1)		518	Initial	7/14/04
532	IND Safety Report	7/28/04			
	Initial report: 2004-BP-05700BP(0)			Initial	
	Follow-up #2: 2004-UK-00597UK(2)		504	Initial	7/2/04
			524	Follow-up#1	7/20/04

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Serial No	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
	Follow-up #1: 2004-BP-04048BP(1) Follow-up #3: 2004-BP-04724BP(3)		457 492 511 525	Initial Initial Follow-up#1 Follow-up#2	6/7/04 6/25/04 7/12/04 7/21/04
533	General Correspondence: Request for FDA Feedback - Resistance Template	7/28/04	N/A N/A	Pre-NDA Meeting FDA email providing an updated template for reporting resistance data in the TPV NDA	6/2/2004 6/29/2004
534	Response to FDA Request for Information – In Vitrol No. 527. BI would like to request feedback on the proposal for design and timing for further study of approved ARV compounds	7/28/04	N/A N/A\ SN 123 SN 213 N/A	Type C Clinical Development Meeting between FDA and BI (FDA minutes issued Nov 30, 2001 BI minutes Early data submission Telefax from FDA	10/5/01 11/30/01 11/7/01 3/18/03 11/18/03
535	Response to FDA Request for Information - Analysis Plan for Step 2 Cohort Analysis	7/28/04	414 N/A	Planned Cohort Analysis Proposal Teleconference	5/14/04

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Ž		Submission	Referenced	of Reference	
	Aanalysis plan describing the planned cohort analysis proposed in BI is actively proceeding with this cohort analysis				
536	Response to FDA Request for Information:	7/28/04			
	General Global Expanded Access Overview		508	Draft EAP protocol	7/1/04
			N/A	Telecon w/FDA requesting EAP overview	7/14/04
537	IND Safety Report	7/29/04	N/A	N/A	N/A
	Initial 2004-BP-05731BP(0) Initial: 2004-DE-03306DE(0)				
538	Information Amendment: Clinical-Safety	7/30/04	N/A	Meeting w/FDA where BIPI	
	Information-July 2004 Quarterly Safety			agreed to submit for a period not	
	Summary	•		less than I yr: Grade 3 & 4 serious	
				cases on a monthly basis, and a	
				safety summary on a quarterly basis. (beginning on 1/19/04)	
539	IND Safety Report	7/30/04	٩Z		
	Initial: 2004-IT-00118IT(0)		NA		
	Follow-up#1: 2004-CN-00253CN(1)		520	Initial Report	7/16/04
540	IND Safety Report	8/3/04			
	2004-SW-00232DB(0)		N/A	N/A	N/A
	2004-BP-05611BP(1)		529	Initial	7/23/04
	2004-BP-04724BP(4)	-	492	Initial	6/25/04

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
			511	Follow-up #1	7/12/04
			525	Follow-up #2	7/21/04
			532	Follow-up #3	7/28/04
	2004-FF-00234FF(1)		387	Initial	4/16/04
541	IND Safety Report	8/4/04			
	Initial 2004-BP-06047BP(0)		N/A	N/A	N/A
	Initial 2004-FF-00440FF(0)		N/A	N/A	N/A
542	IND Safety Report	8/4/04			
	Initial 2004-BP-06030BP(0)		N/A	N/A	N/A
543	IND Safety Report	8/5/04			
	Initial: 2004-BP-05975BP(0)		N/A	N/A	N/A
	Follow-up #1: 2004-BP-05731BP(1)		537	Initial	7/29/04
544	IND Safety Report	8/5/04			
:	Initial: 2004-BP-05944BP(0)	5	N/A	<b>∀</b> /Z	A/N
	Initial: 2004-BP-05859BP(0)		N/A	N/A	N/A
545	Information	8/6/04	N/A	N/A	N/A
	Amendment:Pharmacology/Toxicology				
	(U04-3154, U04-3179, U04-3309, (U04- 3232)				
546	IND Safety Report	8/6/04	544	Initial Report	8/5/04
	Corrected Information for Initial			•	
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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S N		Submission	Referenced	of Reference	
547	IND Safety reports	8/6/04			
	Initial: 2004-BP-06139BP(0)		N/A	N/A	N/A
	Initial: 2004-BP-06185BP(0)		N/A	N/A	N/A
	Follow-up: 2004-BP-05320BP(1)		515	Initial	7/13/04
	Follow-up: 2004-BP-05700BP(1)		532	Initial	7/28/04
548	IND Safety Report Initial: 2004-FF-00101FF(0)	8/9/04			
	Initial: 2004-FF-00256FF(0)				
	Initial: 2004-BP-0621BP(0)				
	Follow-up: 2004-FF-00284FF(2)		422	Initial	5/20/04
	•	*	452	Follow-up #1	6/3/04
	Follow-up: 2004-BP-04375BP(1)		473	Initial	6/18/04
549	Protocol Amendment: 1182.14 (Amendment	8/9/2004	244	Original protocol	8/8/2003
	#3 dated June 22, 2004)		306	Amendments I and 2	2/6/2004
550	IND Safety Report	8/10/04			
	Report 2004-FF-00265FF(0)		548	Wrong CIOMS (2004-FF-00256FF(0) was attached	8/9/04
551	IND Safety Report	8/10/04			
	Initial: 2004-BP-06076BP(0)		N/A	7 Day Fax was sent on 8/2/04	
	Follow-up: 2004-BP-05320BP(1)		515	Initial	7/13/04

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Sellal	Description of Submission	Date of	Serial No.	Description	Date of Reference
2º		Submission	Referenced	of Reference	
552	IND Safety Report Initial: 2004-FF-00459FF(0)	8/11/04	NA	N/A	N/A
553	IND Safety Report 2004-BP-04927BR(0) 2004-BL-00137BL(0)	8/12/2004	N/A	N/A	N/A
554	Information Amendment: Pre-Clinical Toxicology Reports U00-3087 and U00-0389	8/12/2004	N/A	N/A	N/A
No SN	IND Safety Report 7DAY FAX 2004-DE-04275DE(0)	8/12/2004	N/A	N/A	N/A
555	IND Safety Report Initial: 2004-BP-06314BP(0)	8/12/04	N/A	N/A	N/A
256	Information Amendment: Clinical-Safety Information-July 2004 Monthly Report	8/13/04	N/A	N/A	N/A
557	IND Safety Reports Initial: 2004-BP-02018BP(0)	8/13/04	N/A	N/A	N/A
558	IND Safety Report Follow-up: 2003-BP-10942BP(3)	8/16/04	281 323 360	Initial Follow-up #1 Follow-up #2	1/12/04 2/27/04 3/26/04
559	IND Safety Report Initial: 2004-BP-06423BR(0)	8/16/04	N/A	N/A	N/A

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Serial No	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
995	IND Safety Report Initial: 2004-DE-04275DE(0)	8/17/04	N/A	7 Day Fax	8/12/04
561	IND Safety Report Follow-up: 2004-BP-05731BP(2)	8/18/04	537 543	Initial Follow-up #1	7/29/04 8/5/04
562	IND Safety Report Initial: 2004-CN-00285CN(0) Follow-up: 2004-UK-00057UK(4)	8/19/04	N/A 296	N/A Initial	N/A 1/30/04
			313 329	Follow-up #1 Follow-up #2	2/13/04 3/5/04
			341	Follow-up #3	3/15/04
563	General Correspondence: Letter of Authorization for Emergency Treatment Dr. Gregory Storch	8/20/04	N/A	N/A	N/A
564	IA Clinical Reports 1182 .37, 1182.4, 1182.41	8/20/04	N/A	N/A	N/A
565	IND Safety Report Initial: 2004-FF-00068FF(0) Follow-up: 2004-FF-00440FF(1)	8/20/04	N/A 541	N/A Initial	N/A 8/4/04
995	Information Amendment: Clinical (Final Draft Clinical Report: 1182.51)	8/23/04	N/A	N/A	N/A
267	IND Safety Report Initial: 2004-NL-00113NL(0)	8/23/04	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
268	IND Safety Report	8/24/04			
	Follow-up: 2004-FF-00364FF(1)		479	Initial	6/22/04
	Follow-up: 2004-BP-05731BP(3)		537	Initial	7/29/04
			543	Follow-up #1	8/5/04
			561	Follow-up #2	8/18/04
269	IND Safety Report	8/25/04	N/A	N/A	N/A
	Initial: 2004-BP-07028BP(0)	·			
	Initial: 2004-BP-06866MX(0)				
	Initial: 2004-FF-00482FF(0)				
270	Response to FDA Request for Information:	8/25/04	SN 533	Template for reporting resistance	7/28/04
	Resistance Template (FDA 20 August, 2004			data in the TPV NDA	
	Fax) (CD w/datasets provided)				
571	IND Safety Report	8/26/04			
	Follow-up: 2004-BP-06047BP(1)		541	Initial	8/4/04
572	IND Safety Report	8/27/04			
	Follow-up: 2004-DE-04275DE(1)			7 Day Fax	8/12/04
			990	Initial	8/17/04
	Initial: 2003-FF-0011FF(0)		N/A	N/A	N/A
573	IND Safety Report	8/27/04	N/A	N/A	N/A
	Initial: 2003-FF-00618FF(0)				
	Follow-up: 2004-FF-00410FF(2)		518	Initial	7/14/04
			528	Follow-up #1	7/22/04
574	IND Safety Report	8/30/2004			
	Initial Report: 2004-FF-00491FF(0)				
	Follow-up: 2004-BP-05812BP(1)		539	Initial	7/30/2004
	Follow-up: 2004-BP-03787BP(1)		433	Initial	5/26/2004

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>o</sub>		Submission	Referenced	of Reference	
	Follow-up #4: 2004-BP-00114BP(4)		288	Initial	1/20/2004
			308	Follow-up #1	2/9/2004
			359	Follow-up#2	3/24/2004
			382	Follow-up#3	4/14/2004
FAX	IND 7-Day phone fax 2004-BP-07145BP	8/31/2004	N/A	N/A	N/A
575	IND Safety Report Follow-up Report: 2004-	8/31/2004			
	BP-06030BP(1)		542	Initial	8/4/2004
975	IND Safety Report	9/1/2004	N/A	N/A	N/A
	Initial: 2004-CN-00295CN(0)				
FAX	IND 7-Day phone fax 2004-BP-03153BR	9/3/2004	N/A	N/A	N/A
277	Information Amendment: Clinical	9/10/04	N/A	N/A	N/A
	Final Draft Protocol 1182.60				
278	IND Information Amendment	9/15/04			1
	Clinical – Safety Information –		N/A	N/A	N/A
	August 2004 Monthly Report				
579	IND Safety Reports (September 1-15, 2004)	9/15/04	N/A	N/A	N/A
280	Protocol Amendment: New Investigators 182.58	9/23/04	N/A	N/A	N/A
581	Response to FDA Request for Information: Site Information on Trials 1182.12; 1182.48;	9/23/04	N/A	Telefax from FDA requesting a list of all investigators, sites, pt	6/3/04
	1182.14			numbers, discontinuations for	
	The second of th		513	Submission of req listing above.	7/12/04

Serial	Description of Submission	Date of	Serial No.	<b>Description</b>	Date of Reference
0		Submission	Kererced	of Keterence	
585	Information Amendment: Pharmacology/Toxicology	9/23/04	318 N/A	Outline of immunotoxicity Program	2/20/04
	Request for Feedback on Study of				8/30/04
	Immunotoxicity requested by FDA on August 30, 2004			Teleconference discussion of immunotox program	
583	General Correspondence: eSubmission Demonstration	9/27/04	NA	Teleconference confirming TPV electronic submission demo	9/3/04
				scheduled for 10/05/05	
584	General Correspondence: Request for Evaluation of Tradenames	9/30/04	307	Request for Evaluation of Tradenames-VRAY, ELODIUS, ONTINOR	2/6/04
		N/A	N/A	e-mail to Tanima Sinha requesting a delay in the review of TPV tradename pending internal	6/30/04
				decisions on a potential new tradename	
585	IND Safety Reports (September 16-30, 2004)	9/30/04	N/A	N/A	N/A
586	Info amendment: Clinical- Cohort Analysis of Concomitant Medications a report summarizing the results of special	10/1/04	535	Submission of details of proposed cohort analysis.	7/28/04

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No No		Submission	Referenced	of Reference	
	cohort analysis				
		535	414	First described cohort analysis.	5/14/04
587	Information Amendment:	10/7/2004	N/A	N/A	N/A
	Pharmacology/Toxicology and Clinical (ECG Rpt U04-3310 and interim CTR				
588	Protocol Amendment: New Investigators: 1182.17 & 1182.58	10/7/2004	N/A	N/A	N/A
589	Information Amendment: Clinical-Safety Information-September 2004 Monthly Report	10/13/04	N/A	N/A	N/A
400	Promosed Changes in Written Reguest for	10/14/04	186	Dediatric Proposal	11/15/02
	Dedictio Chidise	1071	N/A	Dedictric Weitten Demost	11/12/02
	rediatile Studies		N/A	regiante withen request	1/22/03
			747	Changes to written Kequest	1/23/03
			N/A	Leleconterence	10/02/03
591	IND Safety Reports (October 1-15, 2004)	10/15/04	N/A	N/A	N/A
FAX	IND Safety Report: 7-Day Facsimile 2004-BP-09458BP(0)	10/15/04	N/A	N/A	N/A
FAX	IND Safety Report: 7 -Day Facsimile 2004-BP-09805BP(0)	10/20/04	N/A	N/A	N/A
592	Protocol Amendment: Change in Protocol	10/21/04	078	Original Protocol	2/12/01
	1182.17 Amendment 6		101	Amendment 1	6/25/01
			154	Amendment 2	6/12/02
			222	Amendment 3	4/22/03

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
02		Submission	Keterenced	ot Keterence	
			247	Amendment 4	9/4/03
			404	Amendment 5	5/5/04
593	IND Safety Reports (October 16-30, 2004)	10/28/04	N/A	N/A	N/A
594	Investigators Brochure Version 8	11/1/04	146	Information Amendment - Clinical Indated Investigators Brochure	4/26/02
			210	Version 5	2/25/03
				Information Amendment -	4/16/04
			386	Clinical Updated Investigators	
				Brochure Version 6	
				Clinical	
				Updated Investigators Brochure Version 7	
595	Protocol Amendment New Investigator New Investigators: 1182.17 & 1182.58	11/3/2004	N/A	N/A	N/A
none	IND Safety Report 7 day Fax 2004-CN-00399CN(0)	11/12/2004	11/12/2004	N/A	N/A
969	IND Safety Reports (November 1-15, 2004)	11/15/04	N/A	N/A	N/A
597	General Correspondence LOA for Emergency USE	11/15/04	11/15/04	N/A	N/A
869	Monthly Safety Report October 1-31	11/15/04	11/15/04	N/A	N/A

Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
299	General Correspondence – Emergency Use Authorization – Dr. Meislich	11/16/04	11/16/04	N/A	N/A
009	IND ANNUAL REPORT PROPOSAL TIMELINE	11/18/04	N/A	Meeting between BIPI and Division	6/2/04
			521		7/109/04
				BI meeting minutes	
			N/A	FDA meeting minutes	11/09/04
109	IND Safety Reports (November 16-30, 2004)	11/30/04	N/A	N/A	N/A
602	Information Amendment: Clinical-Safety Information-November 2004 Monthly Report	12/13/04	N/A	N/A	N/A
603	Information Amendment: Clinical: Updated Investigator's Brochure Version 9 Response to Dr. James comments on the IDB	12/13/04	146	Information Amendment - Clinical Updated Investigators Brochure Version 5	4/26/02
-8-	submitted 11/1/04 with updates to the IDB		210	Information Amendment – Clinical Updated Investigators Brochure Version 6	2/25/03
			386	Information Amendment – Clinical Updated Investigators Brochure	4/16/04
				Version 7	

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
			594	Updated Investigators Brochure – Version 8	11/1/04
			N/A	Telefax from Dr. James Commenting on the updated IB submitted on 11/1/04	11/5/04
604	IND Safety Reports (December 1-15, 2004)	12/15/04	N/A	N/A	N/A
909	IND Safety Reports (December 15-30, 2004)	12/30/04	N/A	N/A	N/A
909	Information Amendment: Pharmacology/Toxicology	1/04/05	318	Plans for immunotoxicity testing for TPV	2/20/04
	study summary of unaudited draft results of the immunotoxicity study to NDA 21-814 (Amendment 017; January 4, 2005).		N/A	Teleconference	
209	Information Amendment: CMC	12/14/04	244 306	Pediatric Protocol - 1182.14 Amendments 1 and 2	8/8/03 2/6/04
	provide updated CMC documentation for the oral solution drug product to support on-going and future clinical trials		549	Amendment 3	8/9/04
809	Information Amendment: Clinical-Safety Information-December 2004 Monthly Report	1/10/05	1/13/05	N/A	N/A
FAX	IND 7 Day Fax 2005-BP-00186BP(0)	1/13/05	N/A	N/A	N/A
FAX	IND 7 Day Fax 2005-BP-00303BP(0)	1/14/05	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
609	IND Safety Reports (January 1-15, 2005)	1/14/05	1/14/05	N/A	N/A
610	IND Protocol Amendment New Investigators 1182.17 & 1182.58	1/21/05	1/21/05	N/A	N/A
611	In vitro - virologic interactions	1/21/05	N/A	Pre-NDA meeting held on June 2, 2004.	N/A
	in vitro virologic interactions study (report		763	1. 1. 3	707001
	004-5323).		534	Submission requesting reedback on study design	//28/04
			N/A	FDAs response to 7/28/04 submission	8/12/04
612	IND Safety Reports (January 17-31, 2005)	1/31/05	N/A	N/A	N/A
FAX	IND 7 Day Fax 2005-BP-01108BP(0)	2/1/05	N/A	N/A	N/A
FAX	IND 7 Day Fax 2005-bp-01631BP(0)	2/10/05	N/A/	N/A	N/A
613	Information Amendment: Clinical Version 3 of trial 1182.52 report (U03-3236-03)	2/11/05	N/A	Original NDA	10/21/04
	antiretroviral drug-experienced subjects. Content not changed, just minor formatting.				
FAX	IND 7 Day Fax 2005-BP-01868AU(0)	2/14/05	N/A	N/A	N/A
614	IND Safety Reports (Feb. 1-15, 2005)	2/15/05	N/A	N/A	N/A
615	Monthly Report (Feb. 15, 2005)	2/15/05	N/A	N/A	N/A
616	IND Safety Reports Vital Status cases (February 1-15, 2005)	2/15/05	N/A	N/A	N/A
617	IND ANNUAL REPORT	2/22/05	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
N <sub>0</sub>		Submission	Referenced	of Reference	
618	Information Amendment: Pharm/Tox - mouse carcinogenicity study arm termination – request for FDA feedback	2/23/05	151	Draft Protocol (special protocol assessment)	5/24/02
619	Letter of authorization – Glaxo Smith Kline for protocol GW873140 for IND 65238.	2/25/05	N/A	N/A	N/A
620	IND Safety Reports (February 16-28, 2005)	2/28/05	N/A	N/A	N/A
621	Information Amendment: Pharm/Tox immuno toxicity report U05-3021	3/3/05	SN 318	Overview of Immunotoxicity program	2/20/04
			Teleconference	Discussion of immunotox program	8/30/04
		·	SN 582	Request for feedback on the adequacy of protocol designed for the immunotox study and the planned submission timeline	9/23/04
			A017	Summary of U05-3021	1/4/05
622	Protocol Amendment: Change in Protocol Amendments 7 & 8 to 1182.12 Trial	3/3/05	186	End of Phase II Background Document	11/15/02

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Serial	Description of Submission	Date of Submission	Serial No. Referenced	Description of Reference	Date of Reference
			205	Original Protocol	2/4/03
			214	Amendments land 2	3/19/03
			220	Amendment 3	3/31/03
			230	Amendment 4	6/10/03
			498	Amendment 5 & 6	6/30/04
FAX	IND 7 Day Fax 2005-BP-03684BP(0)	3/10/05	N/A	N/A	N/A
623	Protocol Amendment: Change in Protocol		244	Original protocol	8/8/2003
	1182.14 - Amendments 4 & 5	3/10/05	306	Amendments 1 and 2	2/6/2004
		:	549	Amendment 3	8/9/2004
FAX	IND 7 Day Fax 2005-BP-03525BP(0)	3/11/05	N/A	N/A	N/A
FAX	IND 7 Day Fax 2005-BP-01732BR(0)	3/14/05	N/A	N/A	N/A
624	Monthly Report (March 15, 2005)	3/15/05	N/A	N/A	N/A
625	IND Safety Reports (March 1-15, 2005)	3/15/05	N/A	N/A	N/A
FAX	IND 7 Day Fax 2005-DE-00654DE(0)	3/17/200	N/A	N/A	N/A
979	Information Amendment - Clinical	3/25/05	A029	2 Month Safety Update	2/22/05
	(Microbiology) Tipranavir Resistance Report (U04-3215), to describe the emergent resistance data		A030	Amendment to NDA 21814	2/23/05
FAX	IND 7 Day Fax 2005-IT-0053IT(0)	3/28/05	N/A	N/A	N/A
627	Information Amendment – Pharmacology/Toxicolgy	3/31/05	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
S N		Submission	Referenced	of Reference	-
	U02-3410, U03-3059, U03-3060, U03-3061,				
	U03-3080, U03-3086, U03-3193, U03-3213, 1103-3213, 1103-3213, 1103-3289				
	U03-3578, U03-3583, U04-3006, U04-3028,				
	U04-3030, U04-3084, U04-3101, U04-3102,				
	U04-3110, U04-3112, U04-3132, U04-3183,				
	U04-3233, U04-3234, U04-3307, U04-3371,				
	U04-3372, U04-3406, U04-3529				
628	IND Safety Reports (March 16-31, 2005)	3/31/05	N/A	N/A	N/A
629	Information Amendment:	3/31/05	N/A	N/A	N/A
	Pharmacology/Toxicology				
<u></u>	U04-5551 Ipranavir: Assessment of Testionlar Historiathologic				
	Findings in Toxicity				
630	IND Safety Report: 2005-BL-00074BL(0)	4/1/05	879	Batched Submission	3/31/05
	One pg inadvertently left out of batch				
100	Suominssion Andreas and an arrangement of the party of th	14100	71/1	1/14	2777
031	DESPONSE TO DIT B STITUS	4/4/02	N/A	N/A	N/A
	Drig-Drig Interaction shidy between				
-	Dide-Dide inclaction study octaveni				
	BILK355 and IPV being				
	discontinued				
FAX	IND 7 Day Fax 2005-BP-05595BR(0)	4/12/05	N/A	N/A	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Š		Submission	Referenced	of Reference	
632	IND Safety Reports (Apr 1-15, 2005)	4/15/05	N/A	N/A	N/A
633	General Correspondence(LOA Emergency Use – Dr. Ricaurte	4/18/2005	N/A	N/A	N/A
FAX	IND 7 Day Fax 2005-UK-00633UK(0)	4/22/05	NA	N/A	N/A
634	Protocol Amendment:New Protocol :Change in protocol - 1182.93 bioequivalence study submission of original protocol and Amendment 1	4/27/05	N/A	N/A	N/A
635	Information Amendment CMC information for the drug products to be used in clinical trial 1182.93. This trial is being conducted to establish bioequivalence between 2 batches of Tipranavir Capsules 250 mg maintained under different storage conditions.	4/28/05	N/A	N/A	N/A
636	IND Safety Reports (April 16-29, 2005)	5/03/05	N/A	NA	N/A

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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
No		Submission	Referenced	of Reference	
637	Information Amendment/Pharmacology Toxicology Mouse Carcinogenicity Study	5/3/05	151	Survival mouse carcinogenicity study	5/24/02
	Arm Termination		010	L	30,00,0
			010	Early termination proposal	7/23/03
638	Letter of Authorization in support of drug interaction study of Tipranavir and Reverset in healthy subjects by Incyte corporation	5/4/05	N/A	N/A	N/A
639	Information Amendment/Pharmacology U05-3043, U05-3045 – Antiviral Activity Reports	20/9/5	N/A	N/A	N/A
640	Response to FDA Request for Information Please reference Ms. Sinha's March 16, 2005	5/13/05	e-mail	e-mail query regarding updating the IB and Emergency Access	3/16/05
	e-mail query with comments regarding updating our drug metabolism and drug interaction information in our Investigators			Protocol for drug metabolism and drug interaction information.	
	Brochure (IB) and Emergency Access		teleconference	Agreement to inform investigators	4/27/05
	Protocol- submission of DRAFT letter to investigators			of updated info via letter rather than updating IB and EAP.	
641	IND Safety Reports (May 1 – 15, 2005)	5/18/05	N/A	N/A	N/A
642	General Correspondence - Letter or Authorization for Emergency Use Dr. Mael	5/26/05	N/A	N/A	N/A
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Serial	Description of Submission	Date of	Serial No.	Description	Date of Reference
Ž		Submission	Referenced	of Reference	
643	IND Safety Reports (May 16-31, 2005)	6/3/05	N/A	N/A	N/A
644	IND Safety Reports (June 1-15, 2005)	6/17/05	N/A	N/A	N/A
FAX	7 day fax case 2005-BP-1014BP	6/27/05	N/A	N/A	N/A
FAX	7-day fax case 2005-ES-00183ES	6/27/05	N/A	N/A	N/A
645	IND Safety Reports (June 16-30, 2005)	7/1/05	N/A	N/A	N/A
949	Information Amendment:Clinical:	7/13/05	A080	Post marketing study	6/21/05
	New Protocols:		NDA 21-814	commitments	
	Request for FDA Feedback (Post-Marketing				
	Commitments)				
647	IND Safety Reports (July 1-15, 2005)	7/18/05	N/A	N/A	N/A
648	Information Amendment:CMC	7/18/05	NDA 21814	Cross reference NDA for drug	To date.
	Updated CMC information by cross-reference			substance and drug product	
	to NDA 21814 for drug substance and drug			documentation	
	product documentation				
7 day	7 day facsimile IND Safety Report	7/20/05	7/20/05	N/A	N/A
fax	2005-BP-11727AU(0)				
7 day	7 day facsimile IND safety report:	7/21/05	N/A	N/A	N/A
fax	2005-FF00464FF(0)				

# EXHIBIT G ELIGIBILITY OF PATENT FOR EXTENSION

#### ELIGIBILITY OF U.S. PATENT 5,852,195 FOR EXTENSION

In the opinion of the Applicant, U.S. Patent 5,852,195 is eligible for extension under the provisions of 35 U.S.C. §156, for the reasons which follow.

- (1) The term of this patent has not expired before the submission of this application.
- (2) The term of this patent has never been extended.
- (3) This application for patent term extension is submitted by a registered practitioner on behalf of the record owner of the subject patent, Pharmacia & Upjohn LLC, by virtue of a power of appointment signed by a corporate official submitted simultaneously herewith as Exhibit H.
- (4) The product has been subject to a regulatory review period before commercial marketing or use, pursuant to the provisions of Section 505 of the Federal Food, Drug and Cosmetic Act (21 U.S.C. §355).
- (5) The permission for commercial marketing or use of the product after the regulatory review period is the first permission for commercial marketing or use of the product under the provisions of the Federal Food, Drug and Cosmetic Act.

Applicant believes that the subject patent is entitled to 1278 days of extension.

The claimed length of extension has been calculated in the manner set forth in 37 C.F.R. §1.775, as follows:

Initially, the length of the regulatory review period was determined as set forth in section (c). It is 3115 days, which is the sum of:

- (1) 2931days, the number of days in the period beginning on 13 December 1996, the date the exemption under subsection (I) of section 505 of the Federal Food, Drug and Cosmetic Act for the approved product (IND 51979) became effective, and ending on 21 December 2004, the date the application was initially submitted for such product under section 505(b) of the Federal Food, Drug and Cosmetic Act; and
- (2) 184days, the number of days in the period beginning on 21 December 2004, the date the application (NDA 21-814) was initially submitted for the approved product under section 505 and ending on 22 June 2005, the date such application was approved.

Next, the term of the patent as extended was determined in accordance with subsection (d), by:

(1) subtracting from 3115 days, the number of days calculated above to be in the regulatory review period, 1835 days, which is the sum of the periods set forth in 37 C.F.R. §1.775 (d)(1)(i), (ii) and (iii) as set forth in the table below,

(i) the number of days in the periods of paragraphs (c)(1) and (c))(2) of 37 C.F.R. §1.775 which were on and before the date on which the patent issued	740 days
(ii) the number of days in the periods of paragraphs (c)(1) and (c)(2) of 37 C.F.R. §1.775 during which it is believed it will be determined, under 35 U.S.C. §156(d)(2)(B) by the Secretary of Health and Human Services that the Applicant did not act with due diligence	0 days
(iii) one-half of the number of days remaining in the period defined by paragraph (c)(1) of 37 C.F.R. §1.775 after that period is reduced in accordance with paragraphs (d)(1)(i) and (d)(1)(ii) of 37 C.F.R. §1.775 (ignoring half days for the purposes of subtraction)	1095 days

which calculation yields 1280 days as its result;

- (2) by adding the number of days determined in accordance with 37 C.F.R. §1.775 (d)(1), which is 1280 days, to the original term of the patent as shortened by any terminal disclaimer (which term will expire on 22 December 2015), which calculation yields 24 June 2019 as its result;
- (3) by adding 14 years to 22 June 2005, the date of approval of the application under section 505 of the Federal Food, Drug and Cosmetic Act, which calculation yields 22 June 2019 as its result;
- (4) by comparing 24 June 2019 and 22 June 2019, the dates for the ends of the periods obtained pursuant to 37 C.F.R. §1.775 (d)(2) and (d)(3) with each other and selecting

the earlier date, which comparison yields 22 June 2019 as its result;

- (5) as the original patent was issued after September 24, 1984,
  - (i) by adding five years to 22 December 2015, the original expiration date of the patent or any earlier date set by terminal disclaimer, which calculation yields 22 December 2020; and
  - (ii) by comparing 22 December 2020 and 22 June 2019, the dates obtained pursuant to 37 C.F.R. §1.775 (d)(4) and (d)(5)(i) with each other and selecting the earlier date, which comparison yields 22 June 2019 as its result.

The number of days between 22 December 2015, the original expiration date of the patent, and 22 June 2019, the approval date plus 14 years, is 1278 days, which is the amount of extension claimed.

# EXHIBIT H POWER APPOINTING REGISTERED PRACTITIONER

### IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re

: U.S. Patent 5,852,195

Issued

: December 22, 1998

Inventors

: Romines et al.

For

: PYRANONE COMPOUNDS USEFUL TO TREAT RETROVIRAL

**INFECTIONS** 

Mail Stop Patent Ext.
Director
United States Patent and Trademark Office
P.O. Box 1450
Alexandria, VA 22313-1450

## APPOINTMENT OF AGENT FOR PURPOSE OF

### APPLICATION FOR EXTENSION OF PATENT TERM UNDER 35 U.S.C. § 156

Sir:

Pharmacia & Upjohn Company LLC, a company organized under the laws of Delaware is the assignee and owner of record of U.S. Patent 5,852,195 by virtue of an assignment from each of the individual inventors which was recorded on November 23, 1998 at Reel/Frame 009609/0355.

Pharmacia & Upjohn Company LLC hereby appoints Alan Stempel, Reg. No. 28,991, Thomas Blankinship, Reg. No. 39,909, Anthony P. Bottino, Reg. No. 41,629, Philip I. Datlow, Reg. No. 41,482, Mary-Ellen M. Devlin, Reg. No. 27,928, David A. Dow, Reg. No. 46,124, Michael P. Morris, Reg. No. 34,513, Andrea Small, Reg. No. 54,859 and Timothy X. Witkowski, Reg. No. 40,232, as their attorneys and agents to represent Pharmacia & Upjohn LLC in all matters before the United States Patent and Trademark Office which relate to the filing or prosecution of an application for extension of the term of U.S. Patent 5,852,195 under 35 U.S.C.§ 156.

Respectfully submitted,

Steve T. Zelson C Assistant Secretary



Creation date: 09-09-2005

Indexing Officer: TLAM2 - THY LAM

Team: OIPEBackFileIndexing

Dossier: 08809224

Legal Date: 09-02-2005

No.	Doccode	Number of pages
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Total number of pages: 1

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